



(1) Publication number: 0 386 857 B1

(12)

EUROPEAN PATENT SPECIFICATION

(45) Date of publication of patent specification: 16.09.92 Bulletin 92/38

(51) Int. CI.⁵: C12Q 1/68, // C12N9/12, C12N15/54

(21) Application number: 90201138.6

(2) Date of filing: 24.12.87

54 T7 DNA polymerase.

- 30 Priority: 14.01.87 US 3227 14.12.87 US 132569
- 43 Date of publication of application: 12.09.90 Bulletin 90/37
- (45) Publication of the grant of the patent: 16.09.92 Bulletin 92/38
- (A) Designated Contracting States:
 AT BE CH DE ES FR GB GR IT LI LU NL SE
- (56) References cited:
 PROCEEDINGS OF THE NATL. ACADEMY OF SCIENCES USA, vol. 84,July 1987; S.TABOR et al., pp. 4767-4771
 SCIENCE, vol. 230, 20 December 1985; R.K.SAIKI et al., pp. 1350-1354
 THE JOURNAL OF BIOLOGICAL CHEMISTRY, vol. 262, no. 32, 15 November1987; S.TABOR et al., pp. 15330-15333

- 60 Publication number of the earlier application in accordance with Art. 76 EPC: 0 265 293
- (3) Proprietor: THE PRESIDENT AND FELLOWS OF HARVARD COLLEGE 17 Quincy Street Cambridge, MA 02138 (US)
- (72) Inventor: Tabor, Stanley
 9a Lowell Street
 Cambridge, MA 02138 (US)
 Inventor: Richardson, Charles C.
 78 Chestnut Hill Road
 Chestnut Hill, MA 02167 (US)
- (4) Representative: Moon, Donald Keith et al BREWER & SON Quality House Quality Court Chancery Lane London WC2A 1HT (GB)

86 857 B1

0

Ш

Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid (Art. 99(1) European patent convention).

Description

5

25

40

50

55

This invention relates to the use of bacteriophage T7-type DNA polymerases in a method of amplification of a DNA sequence.

DNA sequencing involves the generation of four populations of single stranded DNA fragments having one defined terminus and one variable terminus. The variable terminus always terminates at a specific given nucleotide base (either guanine (G), adenine (A), thymine (T), or cytosine (C)). The four different sets of fragments are each separated on the basis of their length, on a high resolution polyacrylamide gel; each band on the gel corresponds colinearly to a specific nucleotide in the DNA sequence, thus identifying the positions in the sequence of the given nucleotide base.

Generally there are two methods of DNA sequencing. One method (Maxam and Gilbert sequencing) involves the chemical degradation of isolated DNA fragments, each labelled with a single radiolabel at its defined terminus, each reaction yielding a limited cleavage specifically at one ore more of the four bases (G, A, T or C). The other method (dideoxy sequencing) involves the enzymatic synthesis of a DNA strand. Four separate syntheses are run, each reaction being caused to terminate at a specific base (G, A, T or C) via incorporation of the appropriate chain terminating dideoxynucleotide. The latter method is preferred since the DNA fragments are uniformly labelled (instead of end labelled) and thus the larger DNA fragments contain increasingly more radioactivity. Further, ³⁵S-labelled nucleotides can be used in place of ³²P-labelled nucleotides, resulting in sharper definition; and the reaction products are simple to interpret since each lane corresponds only to either G, A, T or C. The enzyme used for most dideoxy sequencing is the Escherichia coli DNA-polymerase I large fragment ("Klenow"). Another polymerase used is AMV reverse transcriptase.

Summary of the Invention

The invention features a method of amplification of a DNA sequence comprising annealing a first and second primer to opposite strands of a double stranded DNA sequence and incubating the annealed mixture with a processive bacteriophage T7-type DNA polymerase (also refered to hereinafter as "T7-type DNA polymerase") having less than 500 units of exonuclease activity per mg of polymerase, preferably less than I unit, wherein the first and second primers anneal to opposite strands of the DNA sequence; in preferred embodiments the primers have their 3' ends directed toward each other, and the method further comprises, after the incubation step, denaturing the resulting DNA, annealing the first and second primers to the resulting DNA and incubating the annealed mixture with the polymerase; preferably the cycle of denaturing, annealing and incubating is repeated from I0 to 40 times.

This invention provides a bacteriophage T7-type DNA polymerase which is processive, non-discriminating, and can utilize short primers. Further, the polymerase has less than 50% of the exonuclease activity of the naturally associated level of exonuclease activity of said polymerase. These are ideal properties for the above described method.

Other features and advantages of the invention will be apparent from the following description of the preferred embodiments thereof and from the claims.

Description of the Preferred Embodiments

The drawings will first briefly be described.

45 Drawings

Figs. 1-3 are diagrammatic representations of the vectors pTrx-2, mGPI-1, and pGP5-5 respectively;

Fig. 4 is a graphical representation of the selective oxidation of T7 DNA polymerase;

Fig. 5 is a graphical representation of the ability of modified T7 polymerase to synthesize DNA in the presence of etheno-dATP; and

Fig. 6 is a diagrammatic representation of the enzymatic amplification of genomic DNA using modified T7 DNA polymerase.

Fig. 7, 8 and 9 are the nucleotide sequences of pTrx-2, a part of pGP5-5 and mGPI-2 respectively. Fig. 10 is a diagrammatic representation of pGP5-6.

DNA Polymerase

The bacteriophage T7-type DNA polymerase of this invention, which is substantially the same as the one

in cells infected with a T7-type phage (i.e., phage in which the DNA polymerase requires host thioredoxin as a subunit; for example, the T7-type phage is T7, T3, Φ I, Φ II, H, W31, gh-I, Y, Ali22, or SP6, Studier, 95 Virology 70, 1979), is processive, has less than 50% of the exonuclease activity of the naturally associated level of exonuclease activity of said polymerase, does not discriminate against nucleotide analog incorporation, and can utilize small oligonucleotides (such as tetramers, hexamers and octamers) as specific primers. These properties will now be discussed in detail.

Processivity

10

20

25

30

35

45

By processivity is meant that the DNA polymerase is able to continuously incorporate many nucleotides using the same primer-template without dissociating from the template, under conditions normally used for DNA sequencing extension reactions. The degree of processivity varies with different polymerases: some incorporate only a few bases before dissociating (eg. Klenow (about I5 bases), T4 DNA polymerase (about I0 bases), T5 DNA polymerase (about I80 bases) and reverse transcriptase (about 200 bases) (Das et al. J. Biol. Chem. 254:1227 1979; Bambara et al., J. Biol. Chem 253:413, 1978) while others, such as those of the present invention, will remain bound for at least 500 bases and preferably at least 1,000 bases under suitable environmental conditions. Such environmental conditions include having adequate supplies of all four deoxynucleoside triphosphates and an incubation temperature from 10°C-50°C. Processivity is greatly enhanced in the presence of E. coli single stranded binding (ssb), protein.

With processive enzymes termination of a sequencing reaction will occur only at those bases which have incorporated a chain terminating agent, such as a dideoxynucleotide. If the DNA polymerase is non-processive, then artifactual bands will arise during sequencing reactions, at positions corresponding to the nucleotide where the polymerase dissociated. Frequent dissociation creates a background of bands at incorrect positions and obscures the true DNA sequence. This problem is partially corrected by incubating the reaction mixture for a long time (30-60 min) with a high concentration of substrates, which "chase" the artifactual bands up to a high molecular weight at the top of the gel, away from the region where the DNA sequence is read. This is not an ideal solution since a non-processive DNA polymerase has a high probability of dissociating from the template at regions of compact secondary structure, or hairpins. Reinitiation of primer elongation at these sites is inefficient and the usual result is the formation of bands at the same position for all four nucleotides, thus obscuring the DNA sequence.

Analog discrimation

The DNA polymerases of this invention do not discriminate significantly between dideoxy-nucleotide analogs and normal nucleotides. That is, the chance of incorporation of an analog is approximately the same as that of a normal nucleotide or at least incorporates the analog with at least I/10 the efficiency that of a normal analog. The polymerases of this invention also do not discriminate significantly against some other analogs. This is important since, in addition to the four normal deoxynucleoside triphosphates (dGTP, dATP, dTTP and dCTP), sequencing reactions require the incorporation of other types of nucleotide derivatives such as; radioactively- or fluorescently-labelled nucleoside triphosphates, usually for labeling the synthesized strands with ³⁵S, ³²P, or other chemical agents. When a DNA polymerase does not discriminate against analogs the same probability will exist for the incorporation of an analog as for a normal nucleotide. For labelled nucleoside triphosphates this is important in order to efficiently label the synthesized DNA strands using a minimum of radioactivity. Further, lower levels of analogs are required with such enzymes, making the sequencing reaction cheaper than with a discriminating enzyme.

Discriminating polymerases show a different extent of discrimination when they are polymerizing in a processive mode versus when stalled, struggling to synthesize through a secondary structure impediment. At such impediments there will be a variability in the intensity of different radioactive bands on the gel, which may obscure the sequence.

Exonuclease Activity

The DNA polymerase of the invention has less than 50%, preferably less than 1%, and most preferably less than 0.1%, of the normal or naturally associated level of exonuclease activity (amount of activity per polymerase molecule). By normal or naturally associated level is meant the exonuclease activity of unmodified T7-type polymerase. Normally the associated activity is about 5,000 units of exonuclease activity per mg of polymerase, measured as described below by a modification of the procedure of Chase et al. (249 J. Biol. Chem. 4545, 1974). Exonucleases increase the fidelity of DNA synthesis by excising any newly synthesized

bases which are incorrectly basepaired to the template. Such associated exonuclease activities are detrimental to the quality of DNA sequencing reactions. They raise the minimal required concentration of nucleotide precursors which must be added to the reaction since, when the nucleotide concentration falls, the polymerase activity slows to a rate comparable with the exonuclease-activity, resulting in no net DNA synthesis, or even degradation of the synthesized DNA.

More importantly, associated exonuclease activity will cause a DNA polymerase to idle at regions in the template with secondary structure impediments. When a polymerase approaches such a structure its rate of synthesis decreases as it struggles to pass. An associated exonuclease will excise the newly synthesized DNA when the polymerase stalls. As a consequence numerous cycles of synthesis and excision will occur. This may result in the polymerase eventually synthesizing past the hairpin (with no detriment to the quality of the sequencing reaction); or the polymerase may dissociate from the synthesized strand (resulting in an artifactual band at the same position in all four sequencing reactions); or, a chain terminating agent may be incorporated at a high frequency and produce a wide variability in the intensity of different fragments in a sequencing gel. This happens because the frequency of incorporation of a chain terminating agent at any given site increases with the number of opportunities the polymerase has to incorporate the chain terminating nucleotide, and so the DNA polymerase will incorporate a chain-terminating agent at a much higher frequency at sites of idling than at other sites.

An ideal sequencing reaction will produce bands of uniform intensity throughout the gel. This is essential for obtaining the optimal exposure of the X-ray film for every radioactive fragment. If there is variable intensity of radioactive bands, then fainter bands have a chance of going undetected. To obtain uniform radioactive intensity of all fragments, the DNA polymerase should spend the same Interval of time at each position on the DNA, showing no preference for either the addition or removal of nucleotides at any given site. This occurs if the DNA polymerase lacks any associated exonuclease, so that it will have only one opportunity to incorporate a chain terminating nucleotide at each position along the template.

Short primers

20

25

35

The DNA polymerase of the invention is able to utilize primers of 10 bases or less, as well as longer ones. most preferably of 4-20 bases. The ability to utilize short primers offers a number of important advantages to DNA sequencing. The shorter primers are cheaper to buy and easier to synthesize than the usual 15-20-mer primers. They also anneal faster to complementary sites on a DNA template, thus making the sequencing reaction faster. Further, the ability to utilize small (e.g., six or seven base) oligonucleotide primers for DNA sequencing permits strategies not otherwise possible for sequencing long DNA fragments. For example, a kit containing 80 random hexamers could be generated, none of which are complementary to any sites in the cloning vector. Statistically, one of the 80 hexamer sequences will occur an average of every 50 bases along the DNA fragment to be sequenced. The determination of a sequence of 3000 bases would require only five sequencing cycles. First, a "universal" primer (e.g., New England Biolabs #1211, sequence 5' GTAAAACGACGGCCAGT 3') would be used to sequence about 600 bases at one end of the insert. Using the results from this sequencing reaction, a new primer would be picked from the kit homologous to a region near the end of the determined sequence. In the second cycle, the sequence of the next 600 bases would be determined using this primer. Repetition of this process five times would determine the complete sequence of the 3000 bases, without necessitating any subcloning, and without the chemical synthesis of any new oligonucleotide primers. The use of such short primers may be enhanced by including gene 2.5 and 4 protein of T7 in the sequencing reaction.

DNA polymerases of this invention, (i.e., having the above properties) include modified bacteriophage T7-type polymerases. That is the DNA polymerase requires host thioredoxin as a sub-unit, and they are substantially identical to a modified bacteriophage T7 DNA polymerase or to equivalent enzymes isolated from related phage, such as T3, Φ I, Φ II, H, W31, gh-I, Y, All22 and SP6. Each of these enzymes can be modified to have properties similar to those of the modified bacteriophage T7 enzyme. It is possible to isolate the enzyme from phage infected cells directly, but preferably the enzyme is isolated from cells which overproduce it. By substantially identical is meant that the enzyme may have amino acid substitutions which do not affect the overall properties of the enzyme. One example of a particularly desirable amino acid substitution is one in which the natural enzyme is modified to remove any exonuclease activity. This modification may be performed at the genetic or chemical level (see below).

Cloning T7 polymerase

As an example of the invention we shall describe the cloning, overproduction, purification, modification and use of T7 DNA polymerase. This processive enzyme consists of two polypeptides tightly complexed in a one

to one stoichiometry. One is the phage T7-encoded gene 5 protein of 84,000 daltons (Modrich et al. 150 J. Biol. Chem. 5515, 1975), the other is the <u>E. coli</u> encoded thioredoxin, of 12,000 daltons (Tabor et al., J. Biol, Chem. 262:16, 216, 1987). The thioredoxin is an accessory protein and attaches the gene 5 protein (the non-processive actual DNA polymerase) to the primer template. The natural DNA polymerase has a very active 3' to 5' exonuclease associated with it. This activity makes the polymerase useless for DNA sequencing and must be inactivated or modified before the polymerase can be used. This is readily performed, as described below, either chemically, by local oxidation of the exonuclease domain, or genetically, by modifying the coding region of the polymerase gene encoding this activity.

o pTrx-2

In order to clone the <u>trx</u>A (thioredoxin) gene of <u>E. coli</u> wild type <u>E. coli</u> DNA was partially cleaved with <u>Sau</u>3A and the fragments ligated to <u>Bam</u>HI-cleaved T7 DNA isolated from strain T7 ST9 (Tabor et al., in <u>Thioredoxin and Glutaredoxin Systems: Structure and Function</u> (Holmgren et al., eds) pp. 285-300, Raven Press, NY; and Tabor et al., <u>supra</u>). The ligated DNA was transfected into <u>E. coli trx</u>A cells, the mixture plated onto <u>trx</u>A cells, and the resulting T7 plaques picked. Since T7 cannot grow without an active <u>E. coli trx</u>A gene only those phages containing the <u>trx</u>A gene could form plaques. The cloned <u>trx</u>A genes were located on a 470 base pair <u>Hin</u>cli fragment.

In order to overproduce thioreodoxin a plasmid, pTrx-2, was as constructed. Briefly, the 470 base pair Hincll fragment containing the trxA gene was isolated by standard procedure (Maniatis et al., Cloning: ALaboratory Manual, Cold Spring Harbor Labs., Cold Spring Harbor, N.Y.), and ligated to a derivative of pBR322 containing a Ptac promoter (ptac-I2, Amann et al., 25 Gene I67, I983). Referring to Fig. 2, ptac-I2, containing β-lactamase and Col El origin, was cut with Pvull, to yield a fragment of 2290 bp, which was then ligated to two tandem copies of trxA (Hincll fragment) using commercially available linkers (Smal-BamHl polylinker), to form pTrx-2. The complete nucleotide sequence of pTrx-2 is shown in Figure 7. Thioredoxin production is now under the control of the tac promoter, and thus can be specifically induced, e.g. by IPTG (isopropyl β-D-thiogalactoside).

pGP5-5 and mGP1-2

45

Some gene products of T7 are lethal when expressed in <u>E. coli</u>. An expression system was developed to facilitate cloning and expression of, lethal genes, based on the inducible expression of T7 RNA polymerase. Gene 5 protein is lethal in some <u>E. coli</u> strains and an example of such a system is described by Tabor at al. 82 Proc. Nat. Acad. Sci. 1074 (1985) where T7 gene 5 was placed under the control of the Φ10 promoter, and is only expressed when T7 RNA polymerase is present in the cell.

Briefly, pGP5-5 (Fig. 3) was constructed by standard procedures using synthetic <u>Bam</u>HI linkers to join T7 fragment from I4306 (<u>Ndel</u>) to I6869 (<u>Aha</u>III), containing gene 5, to the 560 bp fragment of T7 from 5667 (<u>Hincll</u>) to 6I66 (<u>Fnu</u>4H1) containing both the Φ1.1A and Φ1.1B promoters, which are recognized by T7 RNA polymerase and the 3kb <u>Bam</u>HI-<u>Hincll</u> fragment of pACYC177 (Chang et al., I34 J. Bacteriol. II4I, I978). The nucleotide sequence of the T7 inserts and linkers in shown in Fig. 8. In this plasmid gene 5 is only expressed when T7 RNA polymerase is provided in the cell.

Referring to Fig. 3, T7 RNA polymerase is provided on phage vector mGP1-2. This is similar to pGP1-2 (Tabor et al., id.) except that the fragment of T7 from 3l33 (Haelli) to 5840 (Hinfi), containing T7 RNA polymerase was ligated, using linkers (Bglli and Sall respectively), to BamHI-Sall cut Ml3 mp8, placing the polymerase gene under control of the lac promoter. The complete nucleotide sequence of mGPI-2 is shown in Fig. 9.

Since pGP5-5 and pTrx-2 have different origins of replication (respectively a P15A and a CoIE1 origin) they can be tranformed into one cell simultaneously. pTrx-2 expresses large quantities of thioredoxin in the presence of IPTG. mGP1-2 can coexist in the same cell as these two plasmids and be used to regulate expression of T7-DNA polymerase from pGP5-5, simply by causing production of T7-RNA polymerase by inducing the <u>lac</u> promoter with, e.g., IPTG.

Overproduction of T7 DNA polymerase

There are several potential strategies for overproducing and reconstituting the two gene products of <u>trx</u>A and gene 5. The same cell strains and plasmids can be utilized for all the strategies. In the preferred strategy the two genes are co-overexpressed in the same cell. (This is because gene 5 is susceptible to proteases until thioredoxin is bound to it.) As described in detail below, one procedure is to place the two genes separately on each of two compatible plasmids in the same cell. Alternatively, the two genes could be placed in tandem on the same plasmid. It is important that the T7-gene 5 is placed under the control of a non-leaky inducible pro-

moter, such as Φ 1.1A, Φ 1.1B and Φ 10 of T7, as the synthesis of even small quantities of the two polypeptides together is toxic in most \underline{E} . \underline{coli} cells. By non-leaky is meant that less than 500 molecules of the gene product are produced, per cell generation time, from the gene when the promoter, controlling the gene's expression, is not activated. Preferably the T7 RNA polymerase expression system is used although other expression systems which utilize inducible promoters could also be used. A leaky promoter, e.g., plac, allows more than 500 molecules of protein to be synthesized, even when not induced, thus cells containing lethal genes under the control of such a promoter grow poorly and are not suitable in this invention. It is of course possible to produce these products in cells where they are not lethal, for example, the plac promoter is suitable in such cells.

In a second strategy each gene can be cloned and overexpressed separately. Using this strategy, the cells containing the individually overproduced polypeptides are combined prior to preparing the extracts, at which point the two polypeptides form an active T7 DNA polymerase.

Example 1: Production of T7 DNA polymerase

35

E. coli strain 71.18 (Messing et al., Proc. Nat. Acad. Sci. 74:3642, 1977) is used for preparing stocks of mGP1-71.18 is stored in 50% glycerol at -80°C. and is streaked on a standard minimal media agar plate. A single colony is grown overnight in 25 ml standard M9 media at 37°C, and a single plaque of mGP1-2 is obtained by titering the stock using freshly prepared 7I.I8 cells. The plaque is used to inoculate 10 ml 2X LB (2% Bacto-Tryptone, 1% yeast extract, 0.5% NaCl, 8mM NaOH) containing JMI03 grown to an A₆₉₀=0.5. This culture will provide the phage stock for preparing a large culture of mGP1-2. After 3-12 hours, the 10 ml culture is centrifuged, and the supernatant used to infect the large (2L) culture. For the large culture, 4 X 500 ml 2X LB is inoculated with 4 X 5 ml 71.18 cells grown in M9, and is shaken at 37°C. When the large culture of cells has grown to an A₅₉₀=1.0 (approximately three hours), they are inoculated with 10 ml of supernatant containing the starter lysate of mGP1-2. The infected cells are then grown overnight at 37°C. The next day, the cells are removed by centrifugation, and the supernatant is ready to use for induction of K38/pGP5-5/pTrx-2 (see below). The supernatant can be stored at 4°C for approximately six months, at a titer ~5 X 10^{II} Φ/ml. At this titer, 1 L of phage will infect 12 liters of cells at an A_{690} =5 with a multiplicity of infection of 15. If the titer is low, the mGP1-2 phage can be concentrated from the supernatant by dissolving NaCl (60 gm/liter) and PEG-6000 (65 gm/liter) in the supernatant allowing the mixture to settle at 0°C for 1-72 hours, and then centrifuging (7000 rpm for 20 min). The precipitate, which contains the mGP1-2 phage, is resuspended in approximately 1/20th of the original volume of M9 media.

K38/pGP5-5/pTrx-2 is the \underline{E} . \underline{coli} strain (genotype Hfrc (λ)) containing the two compatible plasmids pGP5-5 and pTrx-2. pGP5-5 plasmid has a P15A origin of replication and expresses the kanamycin (Km) resistance gene. pTrx-2 has a ColEI origin of replication and expresses the ampicillin (Ap) resistance gene. The plasmids are introduced into K38 by standard procedures, selecting KmR and ApR respectively. The cells K38/pGP5-5/pTrx-2 are stored in 50% glycerol at -80°C. Prior to use they are streaked on a plate containing 50µg/ml ampicillin and kanamycin, grown at 37°C overnight, and a single colony grown in 10 ml LB media containing 50μg/ml ampicillin and kanamycin, at 37°C for 4-6 hours. The 10 ml cell culture is used to inoculate 500 ml of LB media containing 50μg/ml ampicillin and kanamycin and shaken at 37°C overnight. The following day, the 500 ml culture is used to inoculate 12 liters of 2X LB-KPO₄ media (2% Bacto-Tryptone, 1% yeast extract, 0.5% NaCl, 20 mM KPO4, 0.2% dextrose, and 0.2% casamino acids, pH 7.4), and grown with aeration in a fermentor at 37°C. When the cells reach an A₆₆₀=5.0 (i.e. logarithmic or stationary phase cells), they are infected with mGP1-2 at a multiplicity of infection of 10, and IPTG is added (final concentration 0.5mM). The IPTG induces production of thioredoxin and the T7 RNA polymerase in mGPI-2, and thence induces production of the cloned DNA polymerase. The cells are grown for an additional 2.5 hours with stirring and aeration, and then harvested. The cell pellet is resuspended in 1.5 L 10% sucrose/20 mM Tris-HCl, pH 8.0/25 mM EDTA and re-spun. Finally, the cell pellet is resuspended in 200 ml 10% sucrose/20 mM Tris-HCl, pH 8/I.0 mM EDTA, and frozen in liquid N₂. From 12 liters of induced cells 70 gm of cell paste are obtained containing approximately 700 mg gene 5 protein and 100 mg thioredoxin.

K38/pTrx-2 (K38 containing pTrx-2 alone) overproduces thioredoxin, and it is added as a "booster" to extracts of K38/pGP5-5/pTrx-2 to insure that thioredoxin is in excess over gene 5 protein at the outset of the purification. The K38/pTrx-2 cells are stored in 50% glycerol at -80°C. Prior to use they are streaked on a plate containing 50 μ g/ml ampicillin, grown at 37°C for 24 hours, and a single colony grown at 37°C overnight in 25 ml LB media containing 50 μ g/ml ampicillin. The 25 ml culture is used to inoculate 2 L of 2X LB media and shaken at 37°C. When the cells reach an A₅₉₀=3.0, the ptac promoter, and thus thioredoxin production, is induced by the addition of IPTG (final concentration 0.5 mM). The cells are grown with shaking for an additional 12-16 hours at 37°C, harvested, resuspended in 600 ml 10% sucrose/20 mM Tris-HCl, pH 8.0/25 mM EDTA, and re-spun. Finally, the cells are resuspended in 40 ml 10% sucrose/20 mM Tris-HCl, pH 8/0.5 mM EDTA, and frozen in

liquid N_2 . From 2L of cells 16 gm of cell paste are obtained containing 150 mg of thioredoxin.

Assays for the polymerase involve the use of single-stranded calf thymus DNA (6mM) as a substrate. This is prepared immediately prior to use by denaturation of double-stranded calf thymus DNA with 50 mM NaOH at 20°C for I5 min., followed by neutralization with HCI. Any purified DNA can be used as a template for the polymerase assay, although preferably it will have a length greater than I,000 bases.

The standard T7 DNA polymerase assay used is a modification of the procedure described by Grippo et al. (246 J. Biol. Chem. 6867, I97I). The standard reaction mix (200 μl final volume) contains 40 mM Tris/HCl pH 7.5, I0 mM MgCl₂, 5 mM dithiothreitol, I00 nmol alkali-denatured calf thymus DNA, 0.3 mM dGTP, dATP, dCTP and [³H]dTTP (20 cpm/pm), 50 μg/ml BSA, and varying amounts of T7 DNA polymerase. Incubation is at 37°C (I0°C-45°C) for 30 min (5 min-60 min). The reaction is stopped by the addition of 3 ml of cold (0°C) I N HCl-0.I M pyrophosphate. Acid-insoluble radioactivity is determined by the procedure of Hinkle et al. (250 J. Biol. Chem. 5523, I974). The DNA is precipitated on ice for I5 min (5 min-12 hr), then precipitated onto glass-fiber filters by filtration. The filters are washed five times with 4 ml of cold (0°C) 0.IM HCl-0.IM pyrophosphate, and twice with cold (0°C) 90% ethanol. After drying, the radioactivity on the filters is counted using a non-aqueous scintillation fluor.

One unit of polymerase activity catalyzes the incorporation of I0 nmol of total nucleotide into an acid-soluble form in 30 min at 37°C, under the conditions given above. Native T7 DNA polymerase and modified T7 DNA polymerase (see below) have the same specific polymerase activity ± 20%, which ranges between 5,000-20,000 units/mg for native and 5,000-50,000 units/mg for modified polymerase) depending upon the preparation, using the standard assay conditions stated above.

T7 DNA polymerase is purified from the above extracts by precipitation and chromatography techniques. An example of such a purification follows.

An extract of frozen cells (200 ml K38/pGP5-5/pTrx-2 and 40 ml K38/pTrx-2) are thawed at 0°C overnight. The cells are combined, and 5 ml of lysozyme (15 mg/ml) and 10 ml of NaCl (5M) are added. After 45 min at 0°C, the cells are placed in a 37°C water bath until their temperature reaches 20°C. The cells are then frozen in liquid N₂. An additional 50 ml of NaCl (5M) is added, and the cells are thawed in a 37°C water bath. After thawing, the cells are gently mixed at 0°C for 60 min. The lysate is centrifuged for one hr at 35,000 rpm in a Beckman 45Ti rotor. The supernatant (250 ml) is fraction I. It contains approximately 700 mg gene 5 protein and 250 mg of thioredoxin (a 2:1 ratio thioredoxin to gene 5 protein).

90 gm of ammonium sulphate is dissolved in fraction I (250 ml) and stirred for 60 min. The suspension is allowed to sit for 60 min, and the resulting precipitate collected by centrifugation at 8000 rpm for 60 min. The precipitate is redissolved in 300 ml of 20 mM Tris-HCl pH 7.5/5 mM 2-mercaptoethanol/0.1 mM EDTA/10% glycerol (Buffer A). This is fraction II.

A column of Whatman DE52 DEAE (12.6 cm² x 18 cm) is prepared and washed with Buffer A. Fraction II is dialyzed overnight against two changes of 1 L of Buffer A each until the conductivity of Fraction II has a conductivity equal to that of Buffer A containing 100 mM NaCl. Dialyzed Fraction II is applied to the column at a flow rate of 100 ml/hr, and washed with 400 ml of Buffer A containing 100 mM NaCl. Proteins are eluted with a 3.5 L gradient from 100 to 400 mM NaCl in Buffer A at a flow rate of 60 ml/hr. Fractions containing T7 DNA polymerase, which elutes at 200 mM NaCl, are pooled. This is fraction III (190 ml).

A column of Whatman P11 phosphocellulose (12.6 cm² x 12 cm) is prepared and washed with 20 mM KPO₄ pH 7.4/5 mM 2-mercaptoethanol/0.1 mM EDTA/10 % glycerol (Buffer B). Fraction III is diluted 2-fold (380 ml) with Buffer B, then applied to the column at a flow rate of 60 ml/hr, and washed with 200 ml of Buffer B containing 100mM KC1. Proteins are eluted with a 1.8 L gradient from 100 to 400 mM KCl in Buffer B at a flow rate of 60 ml/hr. Fractions containing T7 DNA polymerase, which elutes at 300 mM KCl, are pooled. This is fraction IV (370 ml).

A column of DEAE-Sephadex® A-50 (4.9 cm² x 15 cm) is prepared and washed with 20 mM Tris-HCl 7.0/0.1 mM dithiothreitol/0.1 mM EDTA/10% glycerol (Buffer C). Fraction IV is dialyzed against two changes of 1 L Buffer C to a final conductivity equal to that of Buffer C containing 100 mM NaCl. Dialyzed fraction IV is applied to the column at a flow rate of 40 ml/hr, and washed with 150 ml of Buffer C containing 100 mM NaCl. Proteins are eluted with a 1 L gradient from 100 to 300 mM NaCl in Buffer C at a flow rate of 40 ml/hr. Fractions containing T7 DNA polymerase, which elutes at 210 mM NaCl, are pooled. This is fraction V (120 ml).

A column of BioRad HTP hydroxylapatite (4.9 cm² x 15 cm) is prepared and washed with 20 mM KPO₄, pH 7.4/10 mM 2-mercaptoethanol/2 mM Na citrate/10% glycerol (Buffer D). Fraction V is dialyzed against two changes of 500 ml Buffer D each. Dialyzed fraction V is applied to the column at a flow rate of 30 ml/hr, and washed with 100 ml of Buffer D. Proteins are eluted with a 900 ml gradient from 0 to 180 mM KPO₄, pH 7.4 in Buffer D at a flow rate of 30 ml/hr. Fractions containing T7 DNA polymerase, which elutes at 50 mM KPO₄, are pooled. This is fraction VI (130 ml). It contains 270 mg of homogeneous T7 DNA polymerase.

Fraction VI is dialyzed versus 20 mM KPO₄ pH 7.4/0.1 mM dithiothreitol/0.1 mM EDTA/50% glycerol. This

is concentrated fraction VI (~65 ml, 4 mg/ml), and is stored at -20°C.

35

45

55

The isolated T7 polymerase has exonuclease activity associated with it. As stated above this must be inactivated. An example of inactivation by chemical modification follows.

Concentrated fraction VI is dialyzed overnight against 20 mM KPO₄ pH 7.4/0.1 mM dithiothreitol/10% glycerol to remove the EDTA present in the storage buffer. After dialysis, the concentration is adjusted to 2 mg/ml with 20 mM KPO₄ pH 7.4/0.1 mM dithiothreitol/10% glycerol, and 30 ml (2mg/ml) aliquots are placed in 50 ml polypropylene tubes. (At 2 mg/ml, the molar concentration of T7 DNA polymerase is 22 μ M.)

Dithiothreitol (DTT) and ferrous ammonium sulfate (Fe(NH₄)₂(SO₄)₂6H₂O) are prepared fresh immediately before use, and added to a 30 ml aliquot of T7 DNA polymerase, to concentrations of 5 mM DTT (0.6 ml of a 250 mM stock) and 20μ M Fe(NH₄)₂(SO₄)₂6H₂O (0.6 ml of a 1 mM stock). During modification the molar concentrations of T7 DNA polymerase and iron are each approximately 20 μ M, while DTT is in 250X molar excess.

The modification is carried out at 0°C under a saturated oxygen atmosphere as follows. The reaction mixture is placed on ice within a dessicator, the dessicator is purged of air by evacuation and subsequently filled with 100% oxygen. This cycle is repeated three times. The reaction can be performed in air (20% oxygen), but occurs at one third the rate.

The time course of loss of exonuclease activity is shown in Fig. 4. ³H-labeled double-stranded DNA (6 cpm/pmol) was prepared from bacteriophage T7 as described by Richardson (I5 J. Molec. Biol. 49, 1966). ³H-labeled single-stranded T7 DNA was prepared immediately prior to use by denaturation of double-stranded ³H-labeled T7 DNA with 50 mM NaOH at 20°C for I5 min, followed by neutralization with HCl. The standard exonuclease assay used is a modification of the procedure described by Chase et al. (supra). The standard reaction mixture (100 μl final volume) contained 40 mM Tris/HCl pH 7.5, I0 mM MgCl₂, I0 mM dithiothreitol, 60 nmol ³H-labeled single-stranded T7 DNA (6 cpm/pm), and varying amounts of T7 DNA polymerase. ³H-labeled double-stranded T7 DNA can also be used as a substrate. Also, any uniformly radioactively labeled DNA, single- or double-stranded, can be used for the assay. Also, 3′ end labeled singleor double-stranded DNA can be used for the assay. After incubation at 37°C for I5 min, the reaction is stopped by the addition of 30 μl of BSA (I0mg/ml) and 25 μl of TCA (I00% w/v). The assay can be run at I0°C-45°C for I-60 min. The DNA is precipitated on ice for I5 min (I min - I2 hr), then centrifuged at I2,000 g for 30 min (5 min - 3 hr). I00 μl of the supernatant is used to determine the acid-soluble radioactivity by adding it to 400 μl water and 5 ml of aqueous scintillation cocktail.

One unit of exonuclease activity catalyzes the acid solubilization of 10 nmol of total nucleotide in 30 min under the conditions of the assay. Native T7 DNA polymerase has a specific exonuclease activity of 5000 units/mg, using the standard assay conditions stated above. The specific exonuclease activity of the modified T7 DNA polymerase depends upon the extent of chemical modification, but ideally is at least 10-100-fold lower than that of native T7 DNA polymerase, or 500 to 50 or less units/mg using the standard assay conditions stated above. When double stranded substrate is used the exonuclease activity is about 7-fold higher.

Under the conditions outlined, the exonuclease activity decays exponentially, with a half-life of decay of eight hours. Once per day the reaction vessel is mixed to distribute the soluble oxygen, otherwise the reaction will proceed more rapidly at the surface where the concentration of oxygen is higher. Once per day 2.5 mM DTT (0.3 ml of a fresh 250 mM stock to a 30 ml reaction) is added to replenish the oxidized DTT.

After eight hours, the exonuclease activity of T7 DNA polymerase has been reduced 50%, with negligible loss of polymerase activity. The 50% loss may be the result of the complete inactivation of exonuclease activity of half the polymerase molecules, rather than a general reduction of the rate of exonuclease activity in all the molecules. Thus, after an eight hour reaction all the molecules have normal polymerase activity, half the molecules have normal exonuclease activity, while the other half have <0.1% of their original exonuclease activity

When 50% of the molecules are modified (an eight hour reaction), the enzyme is suitable, although suboptimal, for DNA sequencing. For more optimum quality of DNA sequencing, the reaction is allowed to proceed to greater than 99% modification (having less than 50 units of exonuclease activity), which requires four days.

After four days, the reaction mixture is dialyzed against 2 changes of 250 ml of 20 mM KPO₄ pH 7.4/0.1 mM dithiothreitol/0.1 mM EDTA/50% glycerol to remove the iron. The modified T7 DNA polymerase (\sim 4 mg/ml) is stored at -20°C.

The reaction mechanism for chemical modification of T7 DNA polymerase depends upon reactive oxygen species generated by the presence of reduced transition metals such as Fe²⁺ and oxygen. A possible reaction mechanism for the generation of hydroxyl radicals is outlined below:

(1)
$$Fe^{2+} + O_2 \rightarrow Fe^{3+} + O_2$$

(2) $2 O_2 + 2 H^+ \rightarrow H_2 O_2 + O_2$

(3) $Fe^{2+} + H_2O_2 \rightarrow Fe^{3+} + OH + OH^-$

In equation 1, oxidation of the reduced metal ion yields superoxide radical, O2. The superoxide radical can

undergo a dismutation reaction, producing hydrogen peroxide (equation 2). Finally, hydrogen peroxide can react with reduced metal ions to form hydroxyl radicals, OH· (the Fenton reaction, equation 3). The oxidized metal ion is recycled to the reduced form by reducing agents such as dithiothreitol (DTT).

These reactive oxygen species probably inactivate proteins by irreversibly chemically altering specific amino acid residues. Such damage is observed in SDS-PAGE of fragments of gene 5 produced by CNBr or trypsin. Some fragments disappear, high molecular weight cross linking occurs, and some fragments are broken into two smaller fragments.

As previously mentioned, oxygen, a reducing agent (e.g. DTT, 2-mercaptoethanol) and a transition metal (e.g. iron) are essential elements of the modification reaction. The reaction occurs in air, but is stimulated three-fold by use of 100% oxygen. The reaction will occur slowly in the absence of added transition metals due to the presence of trace quantities of transition metals (1-2μM) in most buffer preparations.

As expected, inhibitors of the modification reaction include anaerobic conditions (e.g., N_2) and metal chelators (e.g. EDTA, citrate, nitrilotriacetate). In addition, the enzymes catalase and superoxide dismutase may inhibit the reaction, consistent with the essential role of reactive oxygen species in the generation of modified T7 DNA polymerase.

As an alternative procedure, it is possible to genetically mutate the T7 gene 5 to specifically inactivate the exonuclease domain of the protein. The T7 gene 5 protein purified from such mutants is ideal for use in DNA sequencing without the need to chemically inactivate the exonuclease by oxidation and without the secondary damage that inevitably occurs to the protein during chemical modification.

20

35

Genetically modified T7 DNA polymerase can be isolated by randomly mutagenizing the gene 5 and then screening for those mutants that have lost exonuclease activity, without loss of polymerase activity. Mutagenesis is performed as follows. Single-stranded DNA containing gene 5 (e.g., cloned in pEMBL-8, a plasmid containing an origin for single stranded DNA replication) under the control of a T7 RNA polymerase promoter is prepared by standard procedure, and treated with two different chemical mutagens: hydrazine, which will mutate C's and T's, and formic acid, which will mutate G's and A's. Myers et al. 229 Science 242, 1985. The DNA is mutagenized at a dose which results in an average of one base being altered per plasmid molecule. The single-stranded mutagenized plasmids are then primed with a universal 17-mer primer (see above), and used as templates to synthesize the opposite strands. The synthesized strands contain randomly incorporated bases at positions corresponding to the mutated bases in the templates. The double-stranded mutagenized DNA is then used to transform the strain K38/pGP1-2, which is strain K38 containing the plasmid pGP1-2 (Tabor et al., supra). Upon heat induction this strain expresses T7 RNA polymerase. The transformed cells are plated at 30°C, with approximately 200 colonies per plate.

Screening for cells having T7 DNA polymerase lacking exonuclease activity is based upon the following finding. The 3' to 5' exonuclease of DNA polymerases serves a proofreading function. When bases are misincorporated, the exonuclease will remove the newly incorporated base which is recognized as "abnormal". This is the case for the analog of dATP, etheno-dATP, which is readily incorporated by T7 DNA polymerase in place of dATP. However, in the presence of the 3' to 5' exonuclease of T7 DNA polymerase, it is excised as rapidly as it is incorporated, resulting in no net DNA synthesis. As shown in figure 6, using the alternating copolymer poly d(AT) as a template, native T7 DNA polymerase catalyzes extensive DNA synthesis only in the presence of dATP, and not etheno-dATP. In contrast, modified T7 DNA polymerase, because of its lack of an associated exonuclease, stably incorporates etheno-dATP into DNA at a rate comparable to dATP. Thus, using poly d(AT) as a template, and dTTP and etheno-dATP as precursors, native T7 DNA polymerase is unable to synthesize DNA from this template, while T7 DNA polymerase which has lost its exonuclease activity will be able to use this template to synthesize DNA.

The procedure for lysing and screening large number of colonies is described in Raetz (72 Proc. Nat. Acad. Sci. 2274, 1975). Briefly, the K38/pGP1-2 cells transformed with the mutagenized gene 5-containing plasmids are transferred from the petri dish, where they are present at approximately 200 colonies per plate, to a piece of filter paper ("replica plating"). The filter paper discs are then placed at 42°C for 60 min to induce the T7 RNA polymerase, which in turn expresses the gene 5 protein. Thioredoxin is constitutively produced from the chromosomal gene. Lysozyme is added to the filter paper to lyse the cells. After a freeze thaw step to ensure cell lysis, the filter paper discs are incubated with poly d(AT), [α³²P]dTTP and etheno-dATP at 37°C for 60 min. The filter paper discs are then washed with acid to remove the unincorporated [³²P]dATP. DNA will precipitate on the filter paper in acid, while nucleotides will be soluble. The washed filter paper is then used to expose X-ray film. Colonies which have induced an active T7 DNA polymerase which is deficient in its exonuclease will have incorporated acid-insoluble ³²P, and will be visible by autoradiography. Colonies expressing native T7 DNA polymerase. or expressing a T7 DNA polymerase defective in polymerase activity, will not appear on the autoradiograph.

Colonies which appear positive are recovered from the master petri dish containing the original colonies.

Cells containing each potential positive clone will be induced on a larger scale (one liter) and T7 DNA polymerase purified from each preparation to ascertain the levels of exonuclease associated with each mutant. Those low in exonuclease are appropriate for DNA sequencing.

Directed mutagenesis may also be used to isolate genetic mutants in the exonuclease domain of the T7 gene 5 protein. The following is an example of this procedure.

T7 DNA polymerase with reduced exonuclease activity (modified T7 DNA polymerase) can also be distinguished from native T7 DNA polymerase by its ability to synthesize through regions of secondary structure. Thus, with modified DNA polymerase. DNA synthesis from a labeled primer on a template having secondary structure will result in significantly longer extensions, compared to unmodified or native DNA polymerase. This assay provides a basis for screening for the conversion of small percentages of DNA polymerase molecules to a modified form

The above assay was used to screen for altered T7 DNA polymerase after treatment with a number of chemical reagents. Three reactions resulted in conversion of the enzyme to a modified form. The first is treatment with iron and a reducing agent, as described above. The other two involve treatment of the enzyme with photo-oxidizing dyes, Rose Bengal and methylene blue, in the presence of light. The dyes must be titrated carefully, and even under optimum conditions the specificity of inactivation of exonuclease activity over polymerase activity is low, compared to the high specificity of the iron-induced oxidation. Since these dyes are quite specific for modification of histidine residues, this result strongly implicates histidine residues as an essential species in the exonuclease active site.

There are 23 histidine residues in T7 gene 5 protein. Eight of these residues lie in the amino half of the protein, in the region where, based on the homology with the large fragment of <u>E</u>. <u>coli</u> DNA polymerase I, the exonuclease domain may be located (Ollis et al. Nature 313, 818. 1984). As described below, seven of the eight histidine residues were mutated individually by synthesis of appropriate oligonucleotides, which were then incorporated into gene 5. These correspond to mutants 1, and 6-10 in table 1.

The mutations were constructed by first cloning the T7 gene 5 from pGP5-3 (Tabor et al., J. Biol. Chem. 282, 1987) into the <u>Smal</u> and <u>Hind</u>III sites of the vector M13 mp18, to give mGP5-2. (The vector used and the source of gene 5 are not critical in this procedure.) Single-stranded mGP5-2 DNA was prepared from a strain that incorporates deoxyuracil in place of deoxythymidine (Kunkel, Proc. Natl. Acad. Sci. USA 82, 488, 1985). This procedure provides a strong selection for survival of only the synthesized strand (that containing the mutation) when transfected into wild-type <u>E.coli</u>, since the strand containing uracil will be preferentially degraded.

Mutant oligonucleotides, 15-20 bases in length, were synthesized by standard procedures. Each oligonucleotide was annealed to the template, extended using native T7 DNA polymerase, and ligated using T4 DNA ligase. Covalently closed circular molecules were isolated by agarose gel electrophoresis, run in the presence of 0.5μg/ml ethidium bromide. The resulting purified molecules were then used to transform Ε. coli 71.18. DNA from the resulting plaques was isolated and the relevant region sequenced to confirm each mutation.

The following summarizes the oligonucleotides used to generate genetic mutants in the gene 5 exonuclease. The mutations created are underlined. Amino acid and base pair numbers are taken from Dunn et al., 166 J. Molec. Biol. 477, 1983. The relevant wild type sequences of the region of gene 5 mutated are also shown.

40 Wild type sequence:

```
109 (aa)

Leu Leu Arg Ser Gly Lys Leu Pro Gly Lys Arg Phe Gly Ser His Ala Leu Glu CTT CTG CGT TCC GGC AAG TTG CCC GGA AAA CGC TTT GGG TCT CAC GCT TTG GAG 14677 (T7 bp)

Mutation 1: His 123 -> Ser 123

Primer used: 5' CGC TTT GGA TCC TCC GCT TTG 3'

Mutant sequence:

123

Leu Leu Arg Ser Gly Lys Leu Pro Gly Lys Arg Phe Gly Ser Ser Ala Leu Glu CTT CTG CGT TCC GGC AAG TTG CCC GGA AAA CGC TTT GGA TCC TCC GCT TTG GAG
```

55

50

45

20

25

Mutation 2: Deletion of Ser 122 and His 123 Primer used: 5' GGA AAA CGC TTT GGC GCC TTG GAG GCG 3' 6 base deletion 5 Mutant sequence: 122 123 Leu Leu Arg Ser Gly Lys Leu Pro Gly Lys Arg Phe Gly · · · · Ala Leu Glu CTT CTG CGT TCC GGC AAG TTG CCC GGA AAA CGC TTT GGC --- --- GCC TTG GAG 10 Mutation 3: Ser 122, His 123 → Ala 122, Glu 123 15 Primer used: 5' CGC TTT GGG GCT GAG GCT TTG G 3' Mutant sequence: 122 123 20 Leu Leu Arg Ser Gly Lys Leu Pro Gly Lys Arg Phe Gly Ala Glu Ala Leu Glu CTT CTG CGT TCC GGC AAG TTG CCC GGA AAA CGC TTT GGG GCT GAG GCT TTG GAG Mutation 4: Lys 118, Arg 119 → Glu 118, Glu 119 25 Primer used: 5' 5' G CCC GGG GAA GAG TTT GGG TCT CAC GC 3' Mutant sequence: 118 119 Leu Leu Arg Ser Gly Lys Leu Pro Gly Glu Glu Phe Gly Ser His Ala Leu Glu 30 CTT CTG CGT TCC GGC AAG TTG CCC GGG GAA GAG TTT GGG TCT CAC GCT TTG GAG Mutation 5:. Arg 111, Ser 112, Lys 114 \rightarrow Glu 111, Ala 112, Glu 114 35 Primer used: 5' G GGT CTT CTG GAA GCC GGC GAG TTG CCC GG 3' Mutant sequence: 111 112 Leu Leu Glu Ala Gly Glu Leu Pro Gly Lys Arg Phe Gly Ser His Ala Leu CTT CTG GAA GCC GGC GAG TTG CCC GGA AAA CGC TTT GGG TCT CAC GCT TTG GAG 45 Mutation 6: His 59, His 62 → Ser 59, Ser 62 Primer used: 5' ATT GTG TTC TCC AAC GGA TCC AAG TAT GAC G 3' 50 Wild-type sequence: 59 62 aa: 55 Leu Ile Val Phe His Asn Gly His Lys Tyr Asp Val CTT ATT GTG TTC CAC AAC GGT CAC AAG TAT GAC GTT 55 77 bp: 14515

	Mutant sequence: 59 62
5	Leu Ile Val Phe <u>Ser</u> Asn Gly Ser Lys Tyr Asp Val CTT ATT GTG TTC <u>TC</u> C AAC GG <u>A TC</u> C AAG TAT GAC GTT
	Mutation 7: His 82 → Ser 82
10	Primer used: 5' GAG TTC TCC CTT CCT CG 3'
	Wild-type sequence: aa: 77 82
15	Leu Asn Arg Glu Phe His Leu Pro Arg Glu Asn TTG AAC CGA GAG TTC CAC CTT CCT CGT GAG AAC T7 bp: 14581
	Mutant sequence:
20	Leu Asn Arg Glu Phe <u>Ser</u> Leu Pro Arg Glu Asn TTG AAC CGA GAG TTC <u>TC</u> C DTT CCT CGT GAG AAC
25	Mutation 8: Arg 96, His 99 → Leu 96, Ser 99
	Primer used: 5' CIG TTG ATT ICT TCC AAC CTC 3'
	Wild-type sequence: aa: 93 96 99
30	Val Leu Ser Arg Leu Ile His Ser Asn Leu Lys Asp Thr Ass GTG TTG TCA CGT TTG ATT CAT TCC AAC CTC AAG GAC ACC GAS T7 bp: 14629
35	Mutant sequence:
	96 99 Val Leu Ser Leu Ile Ser Ser Asn Leu Lys Asp Thr As GTG TTG TCA CTG TTG ATT TCT TCC AAC CTC AAG GAC ACC GAT
40	
	Mutation 9: His 190 → Ser 190 Primer used: 5' CT GAC AAA TCT TAC TTC CCT 3'
45	Wild-type sequence:
	aa: 185 190
50	Leu Leu Ser Asp Lys His Tyr Phe Pro Pro Glu CTA CTC TCT GAC AAA CAT TAC TTC CCT GAG T7 bp: 14905
	Mutant sequence:
e <i>e</i>	190 Leu Leu Ser Asp Lys <u>Ser</u> Tyr Phe Pro Pro Glu CTA CTC TCT GAC AAA <u>TC</u> T TAC TTC CCT CAG

	Mutation 10: His 218 → Ser 218
	Primer used: 5' GAC ATT GAA TOT CGT GCT GC 3'
5	Wild-type sequence:
10	aa: 214 Val Asp Ile Glu His Arg Ala Ala Trp Leu Leu Val Asp Ile Glu His Arg CT GCT GCA TGG CTC GTT GAC ATT GAA CAT CGT GCT GCA TGG CTC 17 bp: 14992
	Mutant sequence: 218
15	Val Asp Ile Glu Ser Arg Ala Ala Trp Leu Leu GTT GAC ATT GAA ZCT CGT GCA TGG CTG CTC
	Mutation 11: Deletion of amino acids 118 to 123
20	Primer used: 5' C GGC AAG TTG CCC GGG GCT TTG GAG GCG TGG G 3'
	18 base deletion
25	Wild-type sequence: 109 (aa) 118 122 123 126 Leu Leu Arg Ser Gly Lys Leu Pro Gly Lys Arg Phe Gly Ser His Ala Leu Glu CTT CTG CGT TCC GGC AAG TTG CCC GGA AAA CGC TTT GGG TCT CAC GCT TTG GAG 14677 (T7 bp)
30	
_	Mutant sequence: 117 Leu Leu Arg Ser Gly Lys Leu Pro Gly(6 amino acids)Ala Leu Glu CTT CTG CGT TCC GGC AAG TTG CCC GGG(18 bases)GCT TTG GAG
35	Mutation 12: His 123 → Glu 123
	Primer used: 5' GGG TCT GAG GCT TTG G 3'
40	Primar dised: 3 666 101 D.L.
	Mutant sequence:
45	Leu Leu Arg Ser Gly Lys Leu Pro Gly Lys Arg Phe Gly Ser Glu Ala Leu Glu CTT CTG CGT TCC GGC AAG TTG CCC GGA AAA CGC TTT GGG TCT GAG GCT TTG GAC
50	Mutation 13: (Arg 131, Lys 136, Lys 140, Lys 144, Arg 145 → Glu 131, Glu 136, Glu 140, Glu 144, Glu 145)
55	Primer used: 5' GGT TAT GAG CTC GGC GAG ATG GAG GGT GAA TAC GAA GAC GAC TIT GAG GAA AT

Wild-type sequence:

129(aa) 131 136 140 144 145

Gly Tyr Arg Leu Gly Glu Het Lys Gly Glu Tyr Lys Asp Asp Phe Lys Arg Het Leu Glu Glu
GGT TAT CGC TTA GGC GAG ATG AAG GGT GAA TAC AAA GAC GAC TTT AAG CGT ATG CTT GAA G
14737 (T7 bp)

10 Murant sequence:

129(aa) 131 136 140 144 145
Gly Tyr Glu Leu Gly Glu Het Glu Gly Glu Tyr Glu Asp Asp Phe Glu Glu Het Leu Glu Glu GGT TAT GAG CTC GGC GAG ATG GAG GGT GAA TAC GAA GAC GAC TTT GAG GAA ATG CTT GAA G
14737 (T7 bp)

15

25

Each mutant gene 5 protein was produced by infection of the mutant phage into K38/pGP1-2, as follows. The cells were grown at 30°C to an A₅₉₀=1.0. The temperature was shifted to 42°C for 30 min., to induce T7 RNA polymerase. IPTG was added to 0.5 mM, and a lysate of each phage was added at a moi=10. Infected cells were grown at 37°C for 90 min. The cells were then harvested and extracts prepared by standard procedures for T7 gene 5 protein.

Extracts were partially purified by passage over a phosphocellulose and DEAE A-50 column, and assayed by measuring the polymerase and exonuclease activities directly, as described above. The results are shown in Table 1.

Table 1 SUMMARY OF EXONUCLEASE AND POLYMERASE ACTIVITIES OF T7 GENE 5 MUTANTS

35	Mutant	Exonuclease activity, %	Polymerase activity, &
	[Wild-type]	[100]a	[100] ^b
40	Mutant 1 (His 123 → Ser 123)	10-25	>90
	Mutant 2 (Δ Ser 122, His 123)	0.2-0.4	>90
45	Murant 3 (Ser 122, His 123 → Ala 122, Glu 123)	<2	>90

50

55

Table 1
SUMMARY OF EXONUCLEASE AND POLYMERASE ACTIVITIES OF T7 GENE 5 MUTANTS

10	Mutant	Exonuclease activity, }	Polymerase activity.
10	Mutant 4 (Lys 118, Arg 119 → Glu 118, Glu 119)) <30	>90
15	Mutant 5 (Arg 111, Ser 112, Lys 114 → Giu 111, Ala 112, Glu 114)	>75	> 9 0
	Mutant 6 (His 59, His $62 \rightarrow Ser 59$, Ser 62)	>75	>90
20	Mutant 7 (His 82 \rightarrow Ser 82)	 >75	>90
	Mutant 8 (Arg 96, His 99 \rightarrow Leu 96, Ser 99)	>75	>90
25	Mutant 9 (His 190 → Ser 190)	>75	>90
	Mutant 10 (His 218 → Ser 218)	>75	>90
30	Mutant 11 (Δ Lys 118, Arg 119, Phe 120, Gly 121, Ser 122, His 123)	<0.02	>90
35	Mutant 12 (His 123 → Glu 123)	<30	>90
40	Mutant 13 (Arg 131, Lys 136, Lys 140, Lys 144, A Glu 131, Glu 136, Glu 140, Glu 144, G	rg 145 → lu 145) <30	>90

a. Exonuclease activity was measured on single stranded [3H]T7 DNA. 100% exonuclease activity corresponds to 5,000 units/mg.

Of the seven histidines tested, only one (His 123: mutant 1) has the enzymatic activities characteristic of modified T7 DNA polymerase. T7 gene 5 protein was purified from this mutant using DEAE-cellulose, phosphocellulose, DEAE-Sephadex and hydroxylapatite chromatography. While the polymerase activity was nearly normal (>90% the level of the native enzyme), the exonuclease activity was reduced 4 to 10-fold.

A variant of this mutant was constructed in which both His 123 and Ser 122 were deleted. The gene 5 protein purified from this mutant has a 200-500 fold lower exonuclease activity, again with retention of >90% of the

15

.

50

5

b. Polymerase activity was measured using single-stranded calf thymus DNA. 100% polymerase activity corresponds to 8,000 units/mg.

polymerase activity.

20

25

30

These data strongly suggest that His 123 lies in the active site of the exonuclease domain of T7 gene 5 protein. Furthermore, it is likely that the His 123 is in fact the residue being modified by the oxidation involving iron, oxygen and a reducing agent, since such oxidation has been shown to modify histidine residues in other proteins (Levine, J. Biol. Chem. <u>258</u>: 11823, 1983; and Hodgson et al. Biochemistry <u>14</u>: 5294, 1975). The level of residual exonuclease in mutant II is comparable to the levels obtainable by chemical modification.

Although mutations at His residues are described, mutations at nearby sites or even at distant sites may also produce mutant enzymes suitable in this invention, e.g.. lys and arg (mutants 4 and 15). Similarly, although mutations in some His residues have little effect on exonuclease activity that does not necessarily indicate that mutations near these residues will not affect exonuclease activity.

Mutations which are especially effective include those having deletions of 2 or more amino acids, preferably 6-8, for example, near the His-123 region. Other mutations should reduce exonuclease activity further, or completely.

As an example of the use of these mutant strains the following is illustrative. A pGP5-6 (mutation II)-containing strain has been deposited with the ATCC (see below). The strain is grown as described above and induced as described in Taber et al. J. Biol. Chem. <u>262</u>:16212 (1987). K38/pTrx-2 cells may be added to increase the yield of genetically modified T7 DNA polymerase.

The above noted deposited strain also contains plasmid pGPI-2 which expresses T7 RNA polymerase. This plasmid is described in Tabor et al., Proc. Nat. Acad. Sci. USA 82:1074, 1985 and was deposited with the ATCC on March 22, 1985 and assigned the number 40,175.

Referring to Fig. 10, pGP5-6 includes the following segments:

- 1. EcoRI-SacI-Smal-BamHI polylinker sequence from MI3 mpl0 (2lbp).
- 2. T7 bp 14309 to 16747, that contains the T7 gene 5, with the following modifications:

T7 bp I4703 is changed from an A to a G, creating a Smal site.

T7 bp I4304 to I432I inclusive are deleted (I8 bp).

- 3. Sall-Pstl-HindIII polylinker sequence from Mi3 mp I0 (15 bp)
- 4. pBR322 bp 29 (HindIII site) to pBR322 bp 375 (BamHI site).
- 5. T7 bp 22855 to T7 bp 22927, that contains the T7 RNA Polymerase promoter φl0, with BamHI linkers inserted at each end (82 bp).
- 6. pBR322 bp 375 (BamHI site) to pBR322 bp 436l (EcoRI site).

DNA Sequencing Using Modified T7-type DNA Polymerase

DNA synthesis reactions using modified T7-type DNA polymerase result in chain-terminated fragments of uniform radioactive intensity, throughout the range of several bases to thousands of bases in length. There is virtually no background due to terminations at sites independent of chain terminating agent incorporation (i.e. at pause sites or secondary structure impediments).

Sequencing reactions using modified T7-type DNA polymerase consist of a pulse and chase. By pulse is meant that a short labelled DNA fragment is synthesized; by chase is meant that the short fragment is lengthened until a chain terminating agent is incorporated. The rationale for each step differs from conventional DNA sequencing reactions. In the pulse, the reaction is incubated at 0°C-37°C for 0.5-4 min in the presence of high levels of three nucleotide triphosphates (e.g., dGTP, dCTP and dTTP) and limiting levels of one other labelled, carrier-free, nucleotide triphosphate, e.g., [36S] dATP. Under these conditions the modified polymerase is unable to exhibit its processive character, and a population of radioactive fragments will be synthesized ranging in size from a few bases to several hundred bases. The purpose of the pulse is to radioactively label each primer, incorporating maximal radioactivity while using minimal levels of radioactive nucleotides. In this example, two conditions in the pulse reaction (low temperature, e.g., from 0-20°C, and limiting levels of dATP, e.g., from 0.1µM to lµM) prevent the modified T7-type DNA polymerase from exhibiting its processive character. Other essential environmental components of the mixture will have similar effects, e.g., limiting more than one nucleotide triphosphate or increasing the ionic strength of the reaction. If the primer is already labelled (e.g., by kinasing) no pulse step is required.

In the chase, the reaction is incubated at 45°C for 1-30 min in the presence of high levels (50-500μM) of all four deoxynucleoside triphosphates and limiting levels (1-50μM) of any one of the four chain terminating agents, e.g., dideoxynucleoside triphosphates, such that DNA synthesis is terminated after an average of 50-600 bases. The purpose of the chase is to extend each radioactively labeled primer under conditions of processive DNA synthesis, terminating each extension exclusively at correct sites in four separate reactions using each of the four dideoxynucleoside triphosphates. Two conditions of the chase (high temperature, e.g., from 30-50°C) and high levels (above 50μM) of all four deoxynucleoside triphosphates) allow the modified T7-type

DNA polymerase to exhibit its processive character for tens of thousands of bases; thus the same polymerase molecule will synthesize from the primer-template until a dideoxynucleotide is incorporated. At a chase temperature of 45°C synthesis occurs at >700 nucleotides/sec. Thus, for sequencing reactions the chase is complete in less than a second. <u>ssb</u> increases processivity, for example, when using dITP, or when using low temperatures or high ionic strength, or low levels of triphosphates throughout the sequencing reaction.

Either [α^{35} S]dATP,[α^{32} P]dATP or fluorescently labelled nucleotides can be used in the DNA sequencing reactions with modified T7-type DNA polymerase. If the fluorescent analog is at the 5' end of the primer, then no pulse step is required.

Two components determine the average extensions of the synthesis reactions. First is the length of time of the pulse reaction. Since the pulse is done in the absence of chain terminating agents, the longer the pulse reaction time, the longer the primer extensions. At 0°C the polymerase extensions average 10 nucleotides/sec. Second is the ratio of deoxyribonucleoside triphosphates to chain terminating agents in the chase reaction. A modified T7-type DNA polymerase does not discriminate against the incorporation of these analogs, thus the average length of extension in the chase is four times the ratio of the deoxynucleoside triphosphate concentration to the chain terminating agent concentration in the chase reaction. Thus, in order to shorten the average size of the extensions, the pulse time is shortened, e.g., to 30 sec. and/or the ratio of chain terminating agent to deoxynucleoside triphosphate concentration is raised in the chase reaction. This can be done either by raising the concentration of the chain terminating agent or lowering the concentration of deoxynucleoside triphosphate. To increase the average length of the extensions, the pulse time is increased, e.g., to 3-4 min, and/or the concentration of chain terminating agent is lowered (e.g., from $20\mu M$) In the chase reaction.

Example 2: DNA sequencing using modified T7 DNA polymerase

25

45

The following is an example of a sequencing protocol using dideoxy nucleotides as terminating agents. 9μ of single-stranded M13 DNA (mGP1-2, prepared by standard procedures) at 0.7 mM concentration is mixed with 1 μ l of complementary sequencing primer (standard universal 17-mer, 0.5 pmole primer / μ l) and 2.5 μ l 5X annealing buffer (200 mM Tris-HC₁, pH 7.5, 50 mM MgCl₂) heated to 65°C for 3 min, and slow cooled to room temperature over 30 min. In the pulse reaction, 12.5 μ l of the above annealed mix was mixed with 1 μ l dithiothreitol 0.1 M, 2 μ l of 3 dNTPs (dGTP, dCTP, dTTP) 3 mM each (P.L Biochemicals, in TE), 2.5 μ l [α ²⁵S]dATP, (1500 Ci/mmol, New England Nuclear) and 1 μ l of modified T7 DNA polymerase described in Example 1 (0.4 mg/ml, 2500 units/ml, i.e. 0.4 μ g, 2.5 units) and incubated at 0°C, for 2 min, after vortexing and centrifuging in a microfuge for 1 sec. The time of incubation can vary from 30 sec to 20 min and temperature can vary from 0°C to 37°C. Longer times are used for determining sequences distant from the primer.

4.5 μ l aliquots of the above pulse reaction are added to each of four tubes containing the chase mixes, preheated to 45°C. The four tubes, labeled G, A, T, C, each contain trace amounts of either dideoxy (dd) G, A, T, or C (P-L Biochemicals). The specific chase solutions are given below. Each tube contains 1.5 μ l dATP 1mM, 0.5 μ l 5X annealing buffer (200 mM Tris-HCl, pH 7.5, 50mM MgCl₂), and 1.0 μ l ddNTP 100 μ M (where ddNTP corresponds to ddG,A,T or C in the respective tubes). Each chase reaction is incubated at 45°C (or 30°C-50°C) for 10 min, and then 6 μ l of stop solution (90% formamide, 10mM EDTA, 0.1% xylenecyanol) is added to each tube, and the tube placed on ice. The chase times can vary from 1-30 min.

The sequencing reactions are run on standard, 6% polyacrylamide sequencing gel In 7M urea, at 30 Watts for 6 hours. Prior to running on a gel the reactions are heated to 75°C for 2 min. The gel is fixed in 10% acetic acid, 10% methanol, dried on a gel dryer, and exposed to Kodak OM1 high-contrast autoradiography film overnight.

Example 3: DNA sequencing using limiting concentrations of dNTPs

In this example DNA sequence analysis of mGPI-2 DNA is performed using limiting levels of all four deoxyribonucleoside triphosphates in the pulse reaction. This method has a number of advantages over the protocol in example 2. First, the pulse reaction runs to completion, whereas in the previous protocol it was necessary to interrupt a time course. As a consequence the reactions are easier to run. Second, with this method it is easier to control the extent of the elongations in the pulse, and so the efficiency of labeling of sequences near the primer (the first 50 bases) is increased approximately I0-fold.

 $7 \mu l$ of 0.75 mM single-stranded MI3 DNA (mGPI-2) was mixed with $l \mu l$ of complementary sequencing primer (I7-mer, 0.5 pmole primer/ μ l) and 2 μ l 5X annealing buffer (200 mM Tris-HCl pH 7.5, 50 mM MgCl₂, 250 mM NaCl) heated at 65°C for 2 min, and slowly cooled to room temperature over 30 min. In the pulse reaction I0 μ l of the above annealed mlx was mixed with $l \mu l$ dithiothreitol 0.I M, 2 μ l of 3 dNTPS (dGTP, dCTP, dTTP) I.5 μ M each, 0.5 μ l [α ³⁶S]dATP, (α l0 μ M) (about $l0\mu$ M, 1500 Ci/mmol, New England Nuclear) and 2 μ l modified T7

DNA polymerase (0.1 mg/ml, 1000 units/ml, i.e., $0.2 \mu g$, 2 units) and incubated at 37°C for 5 min. (The temperature and time of incubation can be varied from 20°C-45°C and I-60 min., respectively.)

 $3.5~\mu$ l aliquots of the above pulse reaction were added to each of four tubes containing the chase mixes, which were preheated to 37° C. The four tubes, labeled G, A, T, C, each contain trace amounts of either dideoxy G, A, T, C. The specific chase solutions are given below. Each tube contains $0.5~\mu$ 5X annealing buffer (200 mM Tris-HCl pH 7.5, 50 mM MgCl₂, 250 mM NaCl), I μ l 4dNTPS (dGTP, dATP, dTTP, dCTP) 200 μ M each, and I.0 μ l ddNTP 20 μ M. Each chase reaction is incubated at 37°C for 5 min (or 20°C-45°C and I-60 mln respectively), and then 4 μ l of a stop solution (95% formamide, 20 mM EDTA, 0.05% xylene-cyanol) added to each tube, and the tube placed on ice prior to running on a standard polyacrylamide sequencing gel as described above.

Example 4: Replacement of dGTP with dITP for DNA sequencing

In order to sequence through regions of compression in DNA, i.e., regions having compact secondary structure, it is common to use dITP (Mills et al., 76 Proc. Natl. Acad. Sci. 2232, 1979) or deazaguanosine triphosphate (deaza GTP, Mizusawa et al., 14 Nuc. Acid Res. 1319, 1986). We have found that both analogs function well with T7-type polymerases, especially with dITP in the presence of ssb.. Preferably these reactions are performed with the above described genetically modified T7 polymerase, or the chase reaction is for I-2 min., and/or at 20°C to reduce exonuclease degradation.

Modified T7 DNA polymerase efficiently utilizes dITP or deaza-GTP in place of dGTP. dITP is substituted for dGTP in both the pulse and chase mixes at a concentration two to five times that at which dGTP is used. In the ddG chase mix ddGTP is still used (not ddITP).

The chase reactions using dITP are sensitive to the residual low levels (about 0.0l units) of exonuclease activity. To avoid this problem, the chase reaction times should not exceed 5 min when dITP is used. It is recommended that the four dITP reactions be run in conjunction with, rather than to the exclusion of, the four reactions using dGTP. If both dGTP and dITP are routinely used, the number of required mixes can be minimized by: (I) Leaving dGTP and dITP out of the chase mixes, which means that the four chase mixes can be used for both dGTP and dITP chase reactions. (2) Adding a high concentration of dGTP or dITP (2μ l at 0.5 mM and I-2.5 mM respectively) to the appropriate pulse mix. The two pulse mixes then each contain a low concentration of dCTP,dTTP and [α^{25} S]dATP, and a high concentration of either dGTP or dITP. This modification does not usually adversely effect the quality of the sequencing reactions, and reduces the required number of pulse and chase mixes to run reactions using both dGTP and dITP to six.

The sequencing reaction is as for example 3, except that two of the pulse mixes contain a) 3 dNTP mix for dGTP: $1.5~\mu M$ dCTP,dTTP, and 1~mM dGTP and b) 3 dNTP mix for dITP: $1.5~\mu M$ dCTP,dTTP, and 2 mM dITP. In the chase reaction dGTP is removed from the chase mixes (i.e. the chase mixes contain 30 μM dATP,dTTP and dCTP, and one of the four dideoxynucleotides at 8 μM), and the chase time using dITP does not exceed 5 min.

Deposits

40

35

10

25

Strains K38/pGP5-5/pTrx-2, K38/pTrx-2 and M13 mGP1-2 have been deposited with the ATCC and assigned numbers 67,287, 67,286, and 40,303 respectively. These deposits were made on January 13, 1987. Strain K38/pGPI-2/pGP5-6 was deposited with the ATCC. On December 4, 1987, and assigned the number 67571.

Applicants' and their assignees acknowledge their responsibility to replace these cultures should they die before the end of the term of a patent issued hereon, 5 years after the last request for a culture, or 30 years, whichever is the longer, and its responsibility to notify the depository of the issuance of such a patent, at which time the deposits will be made irrevocably available to the public. Until that time the deposits will be made irrevocably available to the Commissioner of Patents under the terms of 37 CFR Section 1-14 and 35 USC Section 112.

50

55

Other Embodiments

Other embodiments are within the following claims.

The direct enzymatic amplification of genomic DNA sequences has been described, for other polymerases, by Saiki et al., 230 Science 1350, 1985; and Scharf, 233 Science 1076, 1986.

Referring to Fig. 6, enzymatic amplification of a specific DNA region entails the use of two primers which anneal to opposite strands of a double stranded DNA sequence in the region of interest, with their 3' ends directed toward one another (see dark arrows). The actual procedure involves multiple (10-40, preferably 16-20)

cycles of denaturation, annealing, and DNA synthesis. Using this procedure it is possible to amplify a specific region of human genomic DNA over 200,000 times. As a result the specific gene fragment represents about one part in five, rather than the initial one part in a million. This greatly facilitates both the cloning and the direct analysis of genomic DNA. For diagnostic uses, it can speed up the analysis from several weeks to 1-2 days.

Unlike Klenow fragment, where the amplification process is limited to fragments under two hundred bases in length, modified T7-type DNA polymerases should (preferably in conjuction with <u>E</u>. <u>coli</u> DNA binding protein, or ssb, to prevent "snapback formation of single stranded DNA) permit the amplification of DNA fragments thousands of bases in length.

The modified T7-type DNA polymerases are also suitable in standard reaction mixtures: for a) filling in 5' protruding termini of DNA fragments generated by restriction enzyme cleavage; in order to, for example, produce blunt-ended double stranded DNA from a linear DNA molecule having a single stranded region with no 3' protruding termini; b) for labeling the 3' termini of restriction fragments, for mapping mRNA start sites by SI nuclease analysis, or sequencing DNA using the Maxam and Gilbert chemical modification procedure; and c) for in vitro mutagenesis of cloned DNA fragments. For example, a chemically synthesized primer which contains specific mismatched bases is hybridized to a DNA template, and then extended by the modified T7-type DNA polymerase. In this way the mutation becomes permanently incorporated into the synthesized strand. It is advantageous for the polymerase to synthesize from the primer through the entire length of the DNA. This is most efficiently done using a processive DNA polymerase. Alternatively mutagenesis is performed by misincorporation during DNA synthesis (see above). This application is used to mutagenize specific regions of cloned DNA fragments. It is important that the enzyme used lack exonuclease activity. By standard reaction mixture is meant a buffered solution containing the polymerase and any necessary deoxynucleosides, or other compounds.

25 Claims

30

35

45

50

- A method of amplification of a DNA sequence comprising annealing a first and second primer to opposite strands of a double stranded DNA sequence and incubating the annealed mixture with a DNA polymerase characterized in that said polymerase is a processive bacteriophage T7-type DNA polymerase, having less than 50% of the exonuclease activity of the naturally associated level of exonuclease activity of said polymerase.
- 2. A method as claimed in claim 1 further characterized in that said DNA polymerase has less than 500 units of exonuclease activity per mg of polymerase, and in that said first and second primers anneal to opposite strands of said DNA sequence with their 3' ends directed toward each other after annealing, and with the DNA sequence to be amplified located between the two annealed primers.
- 3. A method as claimed in claim 1 or 2 further characterized in that said polymerase possesses sufficient processivity to remain bound to said DNA sequence for at least 500 bases before dissociating.
- 4. A method as claimed in claim 1, 2 or 3 further characterized in that said polymerase has less than 1% of the exonuclease activity naturally associated with said polymerase.
 - 5. A method as claimed in any of the preceding claims in which said polymerase is T7 DNA polymerase.

Patentansprüche

- 1. Verfahren zum Amplifizieren einer DNA-Sequenz, bei dem an gegenüberliegenden Strängen einer zweisträngigen DNA-Sequenz ein erster und ein zweiter Primer angelagert werden und die angelagerte Mischung mit einer DNA-Polymerase inkubiert wird, dadurch gekennzeichnet, daß die Polymerase eine prozessive, T7 DNA-Polymerase von Bakteorophagen T7 ist, die weniger als 50% der Exonucleaseaktivität aufweist, als es dem natürlichen Maß der Excnucleaseaktivität dieser Polymerase entspricht.
- 2. Verfahren nach Anspruch 1, dadurch gekennzeichnet, daß die DNA-Polymerase weniger als 500 Einheiten der Exonucleaseaktivität pro mg Polymerase aufweist und daß der an die gegenüberliegenden Stränge der DNA-Sequenz angelagerte erste und zweite Primer mit ihren 3'-Enden nach dem Anlagern einander entgegengerichtet sind, wobei sich die zu amplifizierende DNA-Sequenz zwischen den beiden

angelagerten Primern befindet.

- Verfahren nach Anspruch 1 oder 2, dadurch gekennzeichnet, daß die Polymerase eine ausreichende Reaktionsfolge besitzt um an der DNA-Sequenz wenigstens 500 Basen lang gebunden bleibt, ehe sie dissoziiert.
- 4. Verfahren nach den Ansprüchen 1, 2 oder 3, dadurch gekennzeichnet, daß die Polymerase weniger als 1% der Exonucleaseaktivität aufweist, die die Polymerase natürlicherweise hat.
- 5. Verfahren nach einem der vorhergehenden Ansprüche, bei dem die Polymerase eine T7 DNA-Polymerase ist.

Revendications

5

15

20

25

30

40

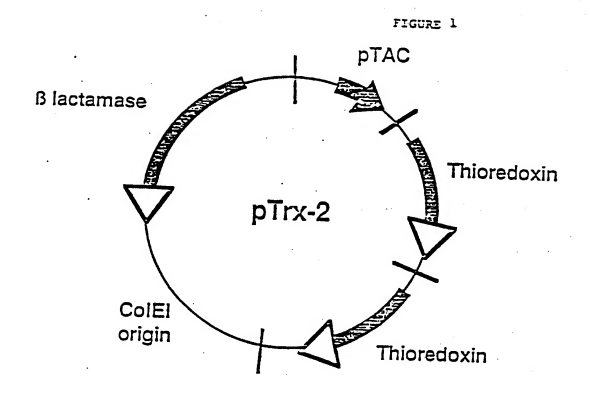
45

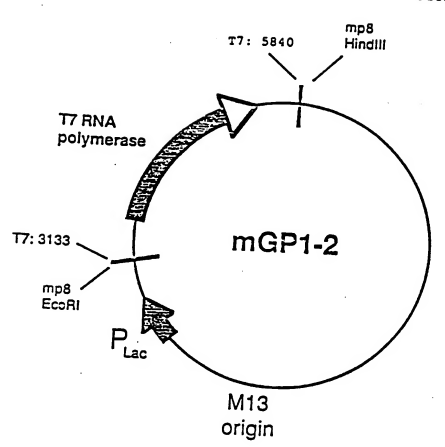
50

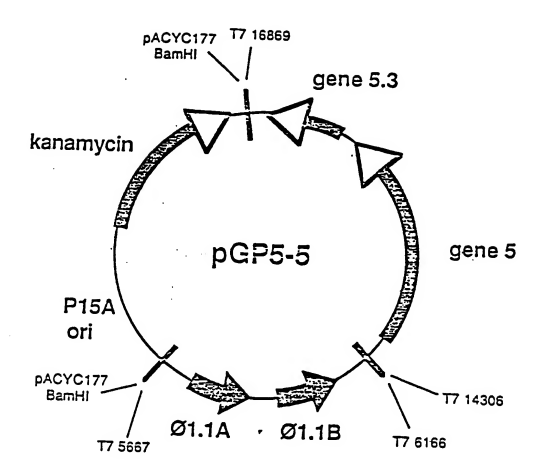
55

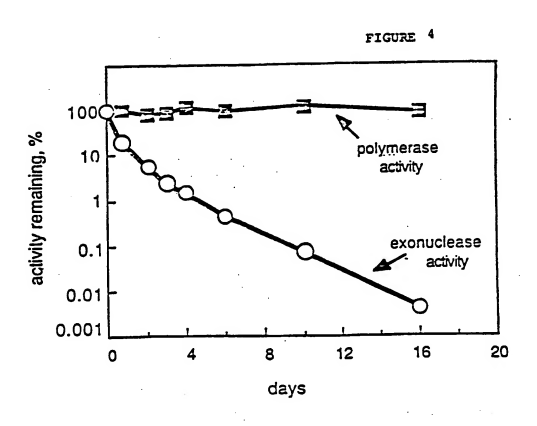
- 1. Procédé d'amplification d'une séquence d'ADN, comprenant l'appariement d'une première et d'une deuxième amorces à des brins opposés d'une séquence d'ADN bicaténaire, et l'incubation du mélange apparié, avec une ADN polymérase, caractérisé en ce que cette polymérase est une ADN polymérase bactériophagique progressive de type T7, ayant moins de 50 % de l'activité d'exonucléase naturelle associée à cette polymérase.
 - 2. Procédé selon la revendication 1, caractérisé en outre en ce que ladite ADN polymérase a une activité d'exonucléase de moins de 500 unités par mg de polymérase, et en ce que les première et deuxième amorces s'apparient à des brins opposés de ladite séquence d'ADN, leurs extrémités 3' étant orientées en direction l'un de l'autre après appariement, et la séquence d'ADN destinée à être amplifiée étant située entre les deux amorces appariées.
 - Procédé selon la revendication 1 ou 2, caractérisé en outre en ce que cette polymérase a une capacité
 de progression suffisante pour rester liée à la séquence d'ADN sur au moins 500 bases avant de se dissocier.
- Procédé selon la revendication 1, 2 ou 3, caractérisé en outre en ce que ladite polymérase a moins de 1 % de l'activité d'exonucléase naturelle associée à cette polymérase.
- Procédé selon l'une quelconque des revendications précédentes, dans lequel ladite polymérase est une
 ADN polymérase T7.

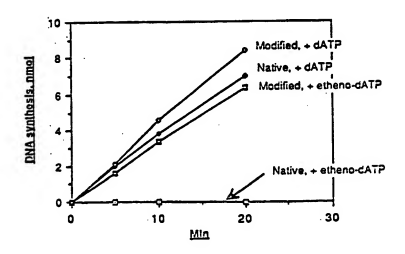
20

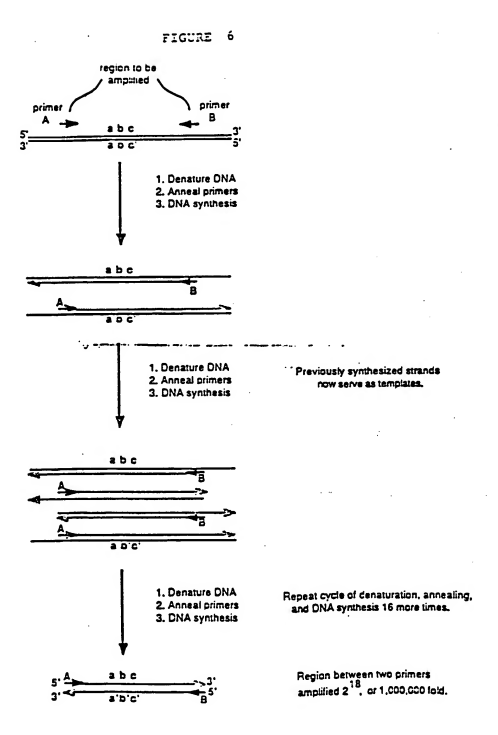












10	20	30	40	50
TTCTTCTCAT	GTTTGACAGC	TTATCATCGA	CTGCACGGTG	CACCAATGCT
60	70	80	90	100
TCTGGCGTCA	GGCAGCCATC	GGAAGCTGTG	GTATGGCTGT	GCAGGTCGTA
110	120	130	140	150
AATCACTGCA	TAATTCGTGT	CGCTCAAGGC	GCACTCCCGT	TCTGGATAAT
160	170		190	200
GTTTTTTGCG	CCGACATCAT	AACGGTTCTG	GCAAATATTC	TGAAATGAGC
210	220	230	240	250
TGTTGACAAT	TAATCATCGG	CTCGTATAAT	GTGTGGAATT	GTGAGCGGAT
260	270		290	300
AACAATTTCA	CACAGGAAAC	AGGGGATCCG	TCAACCTTTA	GTTGGTTAAT
310		330	340	350
GTTACACCAA	CAACGAAACC	AACACGCCAG	GCTTATTCCT	GTGGAGTTAT
360	370	380	390	_. 400
ATATGAGCGA	TAAAATTATT	CACCTGACTG	ACGACAGTTT	TGACACGGAT
410	420	430	440	450
GTACTCAAAG	CGGACGGGGC		GATTTCTGGG	
460			490	500.
CGGTCCGTGC	AAGATGATCG	CCCCGATTCT	GGATGAAATC	GCTGACGAAT

510	520	530	540	550
ATCAGGGCAA	ACTGACCGTT	GCAAAACTGA	ACATCGATCA	AAACCCTGGT
560	570	580	590	600
	AATATGGCAT	CCGTGGTATC	CCGACTCTGC	TGCTGTTCAA
610	620	630	640	650
AAACGGTGAA	GTGGCGGCAA	CCAAAGTGGG	TGCACTGTCT	AAAGGTCAGT
660	670	680	690	700
	CCTCGACGCT	AACCTGGCGT	AAGGGAATTT	CATGTTCGGG
TGAAAGAGTT	720	730	740	750
710		GACGCCCGGC	GTGAGTCATG	CTAACTTAGT
TGCCCCGTCG	CTAAAAACTG	780	790	800
760	770		TTAGTTGGTT	AATGTTACAC
GTTGACGGAT		CCGTCAACCT		850
810	820	830	840	
CAACAACGAA	ACCAACACGC	CAGGCTTATT	CCTGTGGAGT	TATATATGAG
860	870	880	. 890	900
CGATAAAATT	ATTCACCTGA	CTGACGACAG	TTTTGACACG	GATGTACTCA
910	920	930	940	950
AAGCGGACGG	GGCGATCCTC	GTCGATTTCT	GGGCAGAGTG	GTGCGGTCCG
960	970	980	990	1000
		. TCTGGATGAA	ATCGCTGACG	AATATCAGGG
TGCAAGATGA	1020	1030	1040	1050
1010	GTTGCAAAAC	TGAACATCGA		GGTACTGCGC
CAAACTGACC			1090	1100
1060	1070	1080	TGCTGCTGTT	CAAAAACGGT
CGAAATATGG	CATCCGTGGT	ATCCCGACTC		1150
1110	1120	1130	1140	
GAAGTGGCGG	CAACCAAAGT	GGGTGCACTG	TCTAAAGGTC	AGTTGAAAGA
1160	1170	1180	1190	1200
GTTCCTCGAC	GCTAACCTGG	CGTAAGGGAA	TTTCATGTTC	GGGTGCCCCG
1210	1220	1230	1240	1250
TCGCTAAAAA	CTGGACGCCC	GGCGTGAGTC	ATGCTAACTT	AGTGTTGACG
1260	1270	1280	1290	1300
GATCCCCCTG	CCTCGCGCGT	TTCGGTGATG	ACGGTGAAAA	CCTCTGACAC
1310	1320	1330	1340	,1350
	CGGAGACGGT		CTGTAAGCGG	•
ATGCAGCTCC	1370	1380	1390	1400
1360			TGTTGGCGGG	
CAGACAAGCC	CGTCAGGGCG		1440	1450
1410	1420	1430		TGGCTTAACT
CAGCCATGAC	CCAGTCACGT		GAGTGTATAC	
1460	1470	1480	1490	1500
ATGCGGCATC	AGAGCAGATT		TGCACCATAT	
1510	1520		1540	1550
ATACCGCACA	GATGCGTAAG	GAGAAAATAC	CGCATCAGGC	GCTCTTCCGC
1560	1570	1580	1590	
TTCCTCGCTC	ACTGACTCGC	TGCGCTCGGT	CGTTCGGCTG	CGGCGAGCGG
1610			1640	1650
# 3 # C 3 C C T C 3	CTC: 12GCC	GTAATACGGT		ATCAGGGGAT
1660				1700
1000	1070	1000		CCAGGAACCG
1710	1720	1730	1/4U	
		CGTTTTTCCA	AGGCTCCGC	CCCCCTGACG
1760	1770	1780	1790	
AGCATCACAA	AAATCGACGC	: TCAAGTCAGA	GGTGGCGAAA	CCCGACAGGA
1810	1820	1830	1840	1850
CTATAAAGAT	ACCAGGCGTT	TOCCCCTGGA	AGCTCCCTCG	; TGCGCTCTCC

1860	1870	1880	1890	1900
TGTTCCGACC	CTGCCGCTTA		GTCCGCCTTT	
1910	1920	1930	1940	1950
GAAGCGTGGC	GCTTTCTCAA			CAGTTCGGTG
1960	1970	1980	1990	2000
TAGGTCGTTC	GCTCCAAGCT	GGGCTGTGTG	CACGAACCCC	
2010	2020	2030	2040	2050
CGACCGCTGC	GCCTTATCCG	GTAACTATCG		AACCCGGTAA
2060	2070	2080	2090	2100
GACACGACTT	ATCGCCACTG			GATTAGCAGA
2110	2120	2130	2140	2150
GCGAGGTATG	TAGGCGGTGC	TACAGAGTTC	TTGAAGTGGT	GGCCTAACTA
2160	2170	2180	2190	2200
CGGCTACACT	AGAAGGACAG	TATTTGGTAT	CTGCGCTCTG	CTGAAGCCAG
2210	2220	2230	2240	2250
TTACCTTCGG	AAAAAGAGTT	GGTAGCTCTT	GATCCGGCAA	ACAAACCACC
2260	2270	2280	2290	2300
GCTGGTAGCG	GTGGTTTTTT	TGTTTGCAAG	CAGCAGATTA	
2310	2320	2330	2340	2350
AAAAGGATCT	CAAGAAGATC	CTTTGATCTT	TTCTACGGGG	TCTGACGCTC
2360	2370	2380	2390	2400
AGTGGAACGA	AAACTCACGT	TAAGGGATTT	TGGTCATGAG	ATTATCAAAA
2410	2420	2430	2440	2450
AGGATCTTCA	CCTAGATCCT	TTTAAATTAA	AAATGAAGTT	TTAAATCAAT
2460	2470	2480	2490	2500
CTAAAGTATA	TATGAGTAAA	CTTGGTCTGA	CAGTTACCAA	TGCTTAATCA
2510	2520	2530	2540	2550
GTGAGGCACC	TATCTCAGCG	ATCTGTCTAT	TTCGTTCATC	CATAGTTGCC
2560	2570	2580	. 2590	2600
TGACTCCCCG	TCGTGTAGAT	AACTACGATA	CGGGAGGGCT	TACCATCTGG
2610	2620	2.630	2640	2650
CCCCAGTGCT	GCAATGATAC	CGCGAGACCC	ACGCTCACCG	GCTCCAGATT
2660	2670	2680	2690	-2700
TATCAGCAAT	AAACCAGCCA	GCCGGAAGGG	CCGAGCGCAG	AAGTGGTCCT
2710	2720	2730	2740	2750
GCAACTTTAT	CCGCCTCCAT	CCAGTCTATT	AATTGTTGCC	GGGAAGCTAG
2760	2770	2780	2790	2800
AGTAAGTAGT	TCGCCAGTTA	ATAGTTTGCG	CAACGTTGTT	GCCATTGCTG
2810	2820	2830	2840	
CAGGCATCGT	GGTGTCACGC	TCGTCGTTTG	GTATGGCTTC	
2860	2870	2880	2890	
GGTTCCCAAC	GATCAAGGCG	AGTTACATGA	TCCCCCATGT	
2910	2920	2930	2940	
AGCGGTTAGC	TCCTTCGGTC	CTCCGATCGT	TGTCAGAAGT	' AAGTTGGCCG
2960	2970	2980	2990	3000
CAGTGTTATC	ACTCATGGTT	ATGGCAGCAC	TGCATAATTC	TCTTACTGTC
3010	3020	3030	3040	3030
ATGCCATCCG	TAAGATGCTT	TTCTGTGACT	GGTGAGTACT	CAACCAAGTC
30.60	3070	3080	3090	3100
ATTOTGAGAA	TAGTGTATGO	GGCGACCGAG	TTGCTCTTGC	CCGGCGTCAA
3110	3120	3130	3140) .3150
CACCCCATA	TACCGCGCC	CATAGCAGAA	CTTTAAAAGT	GCTCATCATT
3160	3170	3180	3190	3200
CCLLARCETT	CTTCGGGGG	AAAACTCTCA	AGGATCTTAC	CGCTGTTGAG
GGWRRIGG			-	

FIGURE 7 (continued)

3250	3240	3230	3220	3210
TCAGCATCTT	CAACTGATCT	CTCGTGCACC	ATGTAACCCA	ATCCAGTTCG
3300	3290	3280	3270	3260
GCAAAATGCC	AAACAGGAAG	GGGTGAGCAA	CAGCGTTTCT	TTACTTTCAC
3350	3340	3330	3320	3310
TCATACTCTT	TGTTGAATAC	GACACGGAAA	GAATAAGGGC	GCAAAAAAGG
3400	3390	3380	3370	3360
CTCATGAGCG	GGGTTATTGT	GCATTTATCA	TATTATTGAA	CCTTTTTCAA
3450	3440	3430	3420	3410
GGTTCCGCGC	AACAAATAGG	TAGAAAAATA	TGAATGTATT	GATACATATT
3500	3490	3480	3470	3460
TTATTATCAT	TAAGAAACCA	ACCTGACGTC	GAAAAGTGCC	ACATTTCCCC
3550	3540	3530	3520	3510
CGTCTTCAAG	GAGGCCCTTT	GGCGTATCAC	TATAAAAATA	GACATTAACC

AA

10	20	30	40	50
GTTGACACAT	ATGAGTCTTG	TGATGTACTG	GCTGATTTCT	ACGACCAGTT
60	70	80	90	100
CGCTGACCAG			CAAAATGCCA	GCACTTCCGG
110	120	130	140	150
CTAAAGGTAA			TAGAGTCGGA	CTTCGCGTTC
160	170	180	190	200
GCGTAACGCC		GACTCACTAT	AGAGGGACAA	ACTCAAGGTC
210			240	250
	GTGGCCTTTA	TGATTGACCT	TCTTCCGGTT	300
260	270		290	•••
	AACCTTAAGG	TTTAACTTTA	AGACCCTTAA	350
310	320	330	24U	
GAGATTTAAA		ACTAAGAGAG	GACTTTAAGT 390	400
360	370	380	370	
TCGAAAAGAT		TUTAACCGIA	ATGCTCGTGA 440	· 450
410	420	UC# 70077766	CGTGACCGCT	
ACCARAGGTC	GCAAGTTGAA 470	480		500
460	4/0		TATATAGTGG	TAATCTGGCA
TAGCTGGGAG	520		540	. 550
510	72U		ACGATAATCA	
CCGGATCCGG 560	570		590	600
TCAATATGAT	- CG7444C4G7C		ACGCCCTCTT	AGAGAGCGTC'

610	620	630	640	650
ACTAAGTTCC	ACTGCGGGGT	TATCTACGAC	TACTCCACCG	CTGAGTACGT
660	670	680	690	700
AAGCTACCGT	CCGAGTGACT	TCGGTGCGTA	TCTGGATGCG	CTGGAAGCCG
710	720	730	740	750
AGGTTGCACG	AGGCGGTCTT	ATTGTGTTCC	ACAACGGTCA	CAAGTATGAC
760	770	780	790	800
GTTCCTGCAT	TGACCAAACT	GGCAAAGTTG	CAATTGAACC	GAGAGTTCCA
810	820	830	840	850
CCTTCCTCGT		TTGACACCCT	TGTGTTGTCA	CGTTTGATTC
860	870	880	890	900
	CAAGGACACC		TTCTGCGTTC	
ATTCCAACCT		GATATGGGTC		CGGCAAGTTG
910	920	930	940	950
CCCGGAAAAC	GCTTTGGGTC	TCACGCTTTG	GAGGCGTGGG	GTTATCGCTT
. 960	970	980	990	1000
AGGCGAGATG	AAGGGTGAAT	ACAAAGACGA	CTTTAAGCGT	ATGCTTGAAG
1010	1020	1030	1040	1050
AGCAGGGTGA	AGAATACGTT	GACGGAATGG	AGTGGTGGAA	CTTCAACGAA
1060	1070	1080	1090	1100
GAGATGATGG	ACTATAACGT	TCAGGACGTT	GTGGTAACTA	AAGCTCTCCT
1110	1120	1130	1140	1150
TGAGAAGCTA	CTCTCTGACA	AACATTACTT	CCCTCCTGAG	ATTGACTTTA
1160	1170	1180	1190	1200
CGGACGTAGG	ATACACTACG	TTCTGGTCAG	AATCCCTTGA	GGCCGTTGAC
1210	1220	1230	1240	1250
ATTGAACATC	GTGCTGCATG	GCTGCTCGCT	AAACAAGAGC	GCAACGGGTT
1260	1270	1280	1290	1300
CCCGTTTGAC	ACAAAAGCAA	TCGAAGAGTT	GTACGTAGAG	TTAGCTGCTC
1310	1320	1330	1340	1350
GCCGCTCTGA	GTTGCTCCGT	AAATTGACCG	AAACGTTCGG	CTCGTGGTAT
1360	1370		1390	
		1380	CATCCGCGAA	1400
CAGCCTAAAG	GTGGCACTGA	GATGTTCTGC	••••	CAGGTAAGCC
1410	1420	1430	1440	. 1450
ACTACCTAAA	TACCCTCGCA	TTAAGACACC	TAAAGTTGGT	GGTATCTTTA
1460	1470	1480	1490	1500
AGAAGCCTAA	GAACAAGGCA	CAGCGAGAAG	GCCGTGAGCC	TIGCGAACTI
1510	1520	. 1530	1540	1550
GATACCCGCG	AGTACGTTGC	TGGTGCTCCT	TACACCCCAG	TTGAACATGT
1560	1570	1580	1590	1600
TGTGTTTAAC	CCTTCGTCTC	GTGACCACAT	TCAGAAGAAA	CTCCAAGAGG
1610	1620	1630	1640	1650
CTGGGTGGGT	CCCGACCAAG	TACACCGATA	AGGGTGCTCC	TGTGGTGGAC
1660	1670	1680	1690	1700
GATGAGGTAC		ACGTGTAGAT		AGCAAGCCGC
1710	1720	1730	1740	1750
		ACTTGATGAT		
1760	1770	1780	1790	1800
	AGACA22GC2	TGGCTTCGTT	2000000000	CCTCCTASC
· 1810	1820	1830	1840	
		TAATGGAGCA		1850
1860				
	1870	1880	1890	1900
IGCGITCCLA		AAATTCCGGG		
1910	1920	1930	1940	1950
AGCAGTGTCG	CGCTGCTTTT	GGCGCTGAGC	ACCATTTGGA	TGGGATAACT

	1670	1980	1990	2000
1960	1970		GCATCCGGTC	TTGAGCTACG
GGTAAGCCTT	GGGTTCAGGC	TGGCATCGAC	2040	2050
2010	2020	2030	TAACGGCGAG	TACGCTCACG
CTGCTTGGCT	CACTTCATGG	CTCGCTTTGA	2090	2100
2060	2070	2080		
AGATTCTTAA	CGGCGACATC	CACACTAAGA	ACCAGATAGC	TGCTGAACTA
2110	2120	2130	2140	2150
CCTACCCGAG		GACGTTCATC	TATGGGTTCC	TCTATGGTGC
2160	2170	2180	2190	2200
TGGTGATGAG	AAGATTGGAC	AGATTGTTGG	TGCTGGTAAA	GAGCGCGGTA
2210	2220	2230	2240	2250
AGGAACTCAA	GAAGAAATTC	CTTGAGAACA	CCCCGCGAT	TGCAGCACTC
2260	2270	2280	2290	2300
CGCGAGTCTA	TCCAACAGAC	ACTTGTCGAG	TCCTCTCAAT	GGGTAGCTGG
. 2310	2320	2330	2340	2350
TGAGCAACAA	GTCAAGTGGA	AACGCCGCTG	GATTLAAGGT	CTGGATGGTC
2360	2370	2380	2390	2400
GTAAGGTACA	CGTTCGTAGT	CCTCACGCTG	CCTTGAATAC	CCTACTGCAA
2410	2420	2430	2440	2450
TCTGCTGGTG	CTCTCATCTG	CAAACTGTGG	ATTATCAAGA	CCGAAGAGAT
- 2460	2470	2480	2490	2500
GCTCGTAGAG	AAAGGCTTGA	AGCATGGCTG	GGATGGGGAC	TTTGCGTACA
: 2510	2520	2530	2540	2550
TGGCATGGGT	ACATGATGAA	ATCCAAGTAG	GCTGCCGTAC	CGAAGAGATT
2560	2570	2580	2590	2600
GCTCAGGTGG	TCATTGAGAC	CGCACAAGAA	GCGATGCGCT	GGGTTGGAGA
2610	2620	2630	2640	2650
CCACTGGAAC	TTCCGGTGTC	TTCTGGATAC	CGAAGGTAAG	ATGGGTCCTA
2660	2670	2680	. 2690	2700
ATTGGGCGAT	TTGCCACTGA	TACAGGAGGC	TACTCATGAA	CGAAAGACAC
2710	2720	2730	2740	2750
TTAACAGGTG	CTGCTTCTGA	AATGCTAGTA	GCCTACAAAT	TTACCAAAGC
2760	2770	2780	2790	. 2800
TGGGTACACT	GTCTATTACC	CTATGCTGAC	TCAGAGTAAA	GAGGACTTGG
2810	2820	2830	2840	2850
TTGTATGTAA	GGATGGTAAA	TTTAGTAAGG	TTCAGGTTAA	AACAGCCACA
2860	2870	2880	2890	2900
ACGGTTCAAA	CCAACACAGG	AGATGCCAAG	CAGGTTAGGC	TAGGTGGATG
2910	2920	2930	2940	2950
CGGTAGGTCC	GAATATAAGG	ATGGAGACTT	TGACATTCTT	GCGGTTGTGG
2960	2970	2980	2990	3000
TTGACGAAGA	TGTGCTTATT	TTCACATGGG	ACGAAGTAAA	
3010	3020	3030	3040	3050
• : - :	TCGGCAAGAG	AAACAAAGGC	ATAAAACTAT	AGGAGAAATT
TCCATGTGTG	3070	3080	VIWWAY!	HUGHUMAI.
3060		ATTTCCGGAT	С	
ATTATGGCTA	TGACAAAGAA	WILLCOOM		

10	20	30	40	50
ARTGCTACTA		AATTGATGCC	ACCTTTTCAG	CTCGCGCCCC
60	70	80	90	100
AAATGAAAAT	ATAGCTAAAC	AGGTTATTGA	CCATTTGCGA	AATGTATCTA
110	120	130	140	150
ATGGTCAAAC	TAAATCTACT	CGTTCGCAGA	ATTGGGAATC	AACTGTTACA
160	170	180	190	200
TGGAATGAAA	CTTCCAGACA	CCGTACTTTA	GTTGCATATT	TAAAACATGT
210	220	230	240	250
TGAGCTACAG	CACCAGATTC	AGCAATTAAG	CTCTAAGCCA	TCCGCAAAAA
260	270	280	290	300
TGACCTCTTA	TCAAAAGGAG	CAATTAAAGG	TACTCTCTAA	TCCTGACCTG
310	320	330	340	350
TTGGAGTTTG	CTTCCGGTCT	GGTTCGCTTT	GAAGCTCGAA	TTAAAACGCG
360	370	380	390	400
ATATTTGAAG	TCTTTCGGGC	TTCCTCTTAA	TCTTTTTGAT	GCAATCCGCT
410	420	430	440	450
TTGCTTCTGA	CTATAATAGT	CAGGGTAAAG	ACCTGATTTT	TGATTTATGG
4 60	470	480	490	-500
TCATTCTCGT	TTTCTGAACT	GTTTAAAGCA	TTTGAGGGGG	ATTCAATGAA
510	520	530	540	. 550
TATTTATGAC	GATTCCGCAG	TATTGGACGC	TATCCAGTCT	AAACATTTTA
. 560	570	580	590	600
CTATTACCCC	CTCTGGCAAA	ACTICITITG		TCGCTATTTT
610	620	630	640	650
GGTTTTTATC		AAACGAGGGT		TTGCTCTTAC
660	670	680	690	700
TATGCCTCGT	AATTCCTTTT	GGCGTTATGT	ATCTGCATTA	GTTGAATGTG
710	720	730	740	750
GTATTCCTAA	ATCTCAACTG	ATGAATCTTT	CTACCTGTAA	TAATGTTGTT
760	770	780	790	800
CCGTTAGTTC	GTTTTATTAA	CGTAGATTTT	TCTTCCCAAC	GTCCTGACTG
810	820	830	840	850
GTATAATGAG	CCAGTTCTTA	AAATCGCATA	AGGTAATTCA	CAATGATTAA
. 860	870	880	890	900

AGTTGAAATT	AAACCATCTC		TACTACTOGT	TCTGGTGGTT
910	920	930	940	950
CTCGTCAGGG	CAAGCCTTAT	TCACTGAATG	AGCAGCTTTG	TTACGTTGAT
960	970	980	990	1000
TTGGGTAATG			ATTACTCTTG	ATGAAGGTCA
1010	1020	1030	1040	1050
GCCAGCCTAT	GCGCCTGGTC	TGTACACCGT	TCATCTGTCC	TCTTTCAAAG
1060	1070	1080	1090	1100
TTGGTCAGTT	CGGTTCCCTT		GTCTGCGCCT	CGTTCCGGCT
1110	1120	1130	1140	1150
AAGTAACATG	GAGCAGGTCG	CGGATTTCGA	CACAATTTAT	CAGGCGATGA
1160	1170	1180	1190	1200
TACAAATCTC	CGTTGTACTT	TGTTTCGCGC	TTGGTATAAT	CGCTGGGGGT
1210	1220	1230	1240	1250
CAAAGATGAG		TATTCTTTCG	CCTCTTTCGT	TTTAGGTTGG
1260	1270	1280	1290	1300
TGCCTTCGTA	GTGGCATTAC		CGTTTAATGC	AAACTTCCTC
1310	1320	1330	1340	1350
ATGAAAAAGT	CTTTAGTCCT		GTAGCCGTTG	CTACCCTCGT
1360	1370	1380	1390	1400
TCCGATGCTG	TCTTTCGCTG	CTGAGGGTGA	CGATCCCGCA	AAAGCGGCCT
1410	1420	- 1430	1440	1450
TTAACTCCCT	GCAAGCCTCA	GCGACCGAAT	ATATCGGTTA	
1460	1470	1480	1490	1500
ATGGTTGTTG	TCATTGTCGG	CGCAACTATC	GGTATCAAGC	TGTTTAAGAA
1510	1520	1530	1540	1550
ATTCACCTCG	AAAGCAAGCT	GATAAACCGA		GGCTCCTTTT
1560	1570	1580	1590	1600
GGAGCCTTTT	TTTTTGGAGA	TTTTCAACGT		
· 1610	1620	1630	1640	1650
TTCCTTTAGT	TGTTCCTTTC	TATTCTCACT	CCGCTGAAAC	TGTTGAAAGT
1660	1670	1680	. 1690	1700
TGTTTAGCAA	AACCCCATAC	AGAAAATTCA	TTTACTAACG	TCTGGAAAGA
1710	1720	. 1730	1740	1750
CGACAAAACT	TTAGATCGTT	ACGCTAACTA	TGAGGGTTGT	CTGTGGAATG
1760	1770	1780	1790	1800
CTACAGGCGT	TGTAGTTTGT	ACTGGTGACG	AAACTCAGTG	TTACGGTACA
1810	1820	1830	1840	1850
TGGGTTCCTA	TTGGGCTTGC	TATCCCTGAA	AATGAGGGTG	GTGGCTCTGA
1860	1870	1880	1890	1900
GGGTGGCGGT		GCGGTTCTGA	GGGTGGCGGT	ACTABACCTC
1910	1920	1930	1940	1950
	TGATACACCT		ATACTTATAT	
1960				
GACGGCACTT	ATCCGCCTGG	TACTGAGCAA		ATCCTAATCC
2010				2050
##C#C##626	GAGTCTCAGC	CTCTTAATAC	TTTCATGTTT	CAGAATAATA
2060				2100
CCTTCC222	TAGGCAGGGG			CACTGTTACT
2110				2150
2110 2110	20000000000000000000000000000000000000	エコーサンファン		CTGTATCATC
2160				GACTGCGCTT
				2250
2210	2220	2230	2270	2230

	G CTTTAATGA		TTTGTGAATA	TCAAGGCCAA
226				2300
TCGTCTGAC	C TGCCTCAACO	TCCTGTCAAT	GCTGGCGGCG	GCTCTGGTGG
231	0 2320) 2330	2340	
TGGTTCTGG'	T GGCGGCTCTG	AGGGTGGTGG	CTCTGAGGGT	
236		2380		
AGGGTGGCG	G CTCTGAGGGA	GGCGGTTCCG	GTGGTGGCTC	
241				2450
GATTTTGAT:				CTATGACCGA
2460		2480		2500
AAATGCCGA:	GAAAACGCGC	TACAGTCTGA	CGCTAAAGGC	AAACTTGATT
2510	2520	2530		2550
CTGTCGCTAC	TGATTACGGT	GCTGCTATCG	ATGGTTTCAT	TGGTGACGTT
2560		2580		2600
TCCGGCCTTC	CTAATGGTAA	TGGTGCTACT	GGTGATTTTG	CTGGCTCTAA
2610	2620	2630		2650
TTCCCAAATG	GCTCAAGTCG		TAATTCACCT	TTAATGAATA
2660		2680	2690	
ATTTCCGTCA	ATATTTACCT	TCCCTCCCTC	AATCGGTTGA	2700
2710		2730	2740	
	GCGCTGGTAA	ACCATATGLA	2/4V	2750
2760	2770	2780	2790	ATTGTGACAA
	TTCCGTGGTG	TCTTTGCGTT	TCTTTTATAT	2800
2810		2830	2840	GTTGCCACCT
TTATGTATGT			TACTGCGTAA	2850
2860		2880	2890	
TAATCATGCC	AGTTCTTTTG			2900
2910	2920	2930	TATTATTGCG 2940	TTTCCTCGGT
TTCCTTCTGG	TAACTTTGTT			2950
2960	2970	2980	CTTACTTTTC	TTAAAAAGGG
CTTCGGTAAG	ATAGCTATTG	CTATTTCATT	2990	3000
3010	3020	3030	GTTTCTTGCT	CTTATTATTG
	AATTCTTGTG		3040	3050
3060	3070	GGTTATCTCT		•
CCCTCTGACT	TTGTTCAGGG	3080	3090	3100
3110	3120		ATTCTCCCGT	CTAATGCGCT
TCCCTGTTTT	TATGTTATTC	3130	3140	3150
3160		TCTCTGTAAA		TTCATTTTTG
	3170	3180	3190	3200
3210	AAAAATCGTT			ATAATATGGC
TGTTTATTTT	3220	3230	3240	3250
3260	GIAACIGGCA	AATTAGGCTC	TGGAAAGACG	CTCGTTAGCG
TTGGTAAGAT	3270	3280	3290	3300
	TCAGGATAAA	ATTGTAGCTG	GGTGCAAAAT	AGCAACTAAT
3310	3220	4.4.30	2210	
CTIGATTIAA	GGCTTCAAAA	CCTCCCGCAA	GTCGGGAGGT	TCGCTAAAAC
2200	3370	3.350	3300	
GCCTCGCGTT	CTTAGAATAC	CGGATAAGCC	TICTATATCT	GATTTGCTTG
3410	2920	4440	3/40	
CTATTGGGCG	CGGTAATGAT	TCCTACGATG	AAAATAAAA	CGGCTTGCTT
3400	34/0	4490	3400	
GTTCTCGATG	AGTGCGGTAC	TIGGITTAAT	ACCCGTTCTT	GGAATGATAA
3310	352C	マミマハ	3540	200
GGAAAGACAG	CCGATTATTG .	ATTGGTTTCT	ACATGCTCGT	TETRETTAGA
3560	3570	3580	3590	3600
			3230	3 000

GGGATATTAI			CTATTGTTGA	TAAACAGGCG
3610				3650
CGTTCTGCAT	' TAGCTGAACA	TGTTGTTTAT	TGTCGTCGTC	TGGACAGAAT
3660	3670	3680	3690	3700
TACTTTACCT	TTTGTCGGTA	CTTTATATTC	TCTTATTACT	
3710			3740	3750
TGCCTCTGCC			TTAAATATGG	
3760				
TTAAGCCCTA	• • • •		3790	3800
			ACTGGTAAGA	ATTTGTATAA
3810	3820		3840	3850
CGCATATGAT			TAATTATGAT	TCCGGTGTTT
3860	3870		3890	3900
ATTCTTATTT	AACGCCTTAT	TTATCACACG	GTCGGTATTT	CAAACCATTA
3910	3920	3930	3940	3950
AATTTAGGTC	AGAAGATGAA	ATTAACTAAA	ATATATTTGA	AAAAGTTTTC
3960	3970		3990	4000
TCGCGTTCTT	TGTCTTGCGA			
4010	4020		ATCAGCATTT	ACATATAGTT
	ACCTAAGCCG	4030	4040	4050
			AGGTAGTCTC	TCAGACCTAT
4060	4070	4080	4090	4100
	AATTCACTAT		CAGCGTCTTA	ATCTAAGCTA
4110	4120	4130	4140	4150
TCGCTATGTT	. TTCAAGGATT	CTAAGGGAAA	ATTAATTAAT	AGCGACGATT
4160	4170	4180	4190	4200
TACAGAAGCA	AGGTTATTCA	CTCACATATA	TTGATTTATG	TACTGTTTCC
4210	4220	4230	4240	4250
ATTAAAAAAG	GTAATTCAAA	TGAAATTGTT	AAATGTAATT	AATTTTGTTT
4260	4270	4280	4290	4300
TCTTGATGTT		TCTTCTTTTG	CTCAGGTAAT	TGAAATGAAT
4310	4320	4330	4340	
AATTCGCCTC	TGCGCGATTT	TGTAACTTGG		4350
4360	4370		TATTCAAAGC	AATCAGGCGA
ATCCGTTATT	GTTTCTCCCG	4380	4390	4400
		ATGTAAAAGG	TACTGTTACT	GTATATICAT
4410	4420	. 4430	4440	4450
CTGACGTTAA	ACTTGAAAAT	CTACGCAATT	TCTTTATTTC	TGTTTTACGT
4460	4470	4480	4490	4500
GCTAATAATT	TIGATATGGT	TGGTTCAATT	CCTTCCATAA	TTCAGAAGTA
4510	4520	4530	4540	4550
TAATCCAAAC	AATCAGGTAT	ATATTGATGA	ATTGCCATCA	TCTGATAATC
4560	4570	4580	4590	4600
AGGAATATGA		GCTCCTTCTG	منائبنتاكك	
4610	4620	4630		TGTTCCGCAA
LATGATAATG	TTECTTES	UCOF	4640	4650
4660	11701070	TTTAAAATT		GGGCAAAGGA
000F	4670	4680	4690	4700
TITAMIACGA	GITGTCGAAT	TGTTTGTAAA	GTCTAATACT	TCTAAATCCT
4710	4720	4730	4740	4750
CAAATGTATT	ATCTATTGAC	GGCTCTAATC	TATTAGTTGT	TAGTGCACCT
4/60	4770	4780	4790	. 4000
AAAGATATTT	TAGATAACCT	TCCTCAATTC	CTTTCTACTS	TOSTTTTCCC
4810	4820	4830	4840	4950
	ATATTGATTG	AGGGTTTGAT	TUTTE STOTE	0000
4860	4870	4880		
		400V	4890	4900
ADALLOULA	11140111	GCTGCTGGCT		
4910	4920	4930	4940	4950

GGCGGTGTTA	ATACTGACCG	CCTCACCTCT		CIGCIGGIGG
4960	4970	4980	4990	5000
TTCGTTCGGT	ATTTTTAATG	GCGATGTTTT	AGGGCTATCA	GTTCGCGCAT
5010	5020	5030	5040	5050
TAAAGACTAA	TAGCCATTCA	AAAATATTGT	CTGTGCCACG	TATTCTTACG
5060	5070	5080	·5090	5100
CTTTCAGGTC	AGAAGGGTTC		GGCCAGAATG	TCCCTTTTAT
5110	5120	5130	5140	5150
TACTGGTCGT			TGTAAATAAT	CCATTTCAGA
5160	5170	5180	5190	5200
CGATTGAGCG		GGTATTTCCA		TCCTGTTGCA
5210	5220	5230	5240	5250
			ACCAGCAAGG	
ATGGCTGGCG	GTAATATTGT	TCTGGATATT 5280	5290	CCGATAGTTT
5260	5270			5300
GAGTTCTTCT			TACTAATCAA	
5310	5320	5330	5340	5350
CTACAACGGT	TAATTTGCGT	GATGGACAGA	CTCTTTTACT	CGGTGGCCTC
5360	5370	5380	5390	5400
ACTGATTATA		TCAAGATTCT	GGCGTACCGT	TCCTGTCTAA
5410	5420	5430	5440	5450
AATCCCTTTA	ATCGGCCTCC	TGTTTAGCTC	CCGCTCTGAT	TCCAACGAGG
5460	5470	5480	5490	5500
AAAGCACGTT	ATACGTGCTC	GTCAAAGCAA	CCATAGTACG	CGCCCTGTAG
5510	5520	5530	5540	5550
CGGCGCATTA	AGCGCGGCGG	GTGTGGTGGT	TACGCGCAGC	GTGACCGCTA
5560	5570	5580	5590	5600
CACTTGCCAG	CGCCCTAGCG	CCCGCTCCTT	TCGCTTTCTT	CCCTTCCTTT
5610	5620	5630	5640	5650
CTCGCCACGT		TCCCCGTCAA		GGGGGCTCCC
5660	5670	5680	5690	5700
TTTAGGGTTC	CGATTTAGTG	CTTTACGGCA		AAAAAACTTG
5710	5720	5730	5740	5750
ATTTGGGTGA		AGTGGGCCAT		GACGGITTTT
5760	5770	. 5780	5790	5800
CGCCCTTTGA		CACGTTCTTT	AATAGTGGAC	TCTTGTTCCA
5810	5820		5840	5850
		5830		
AACTGGAACA			CTATTCTTTT	GATTTATAAG
5860	5870	5880	5890	5900
GGATTTTGCC		CCACCATCAA		CGCCTGCTGG
5910	5920	5930	5940	5950
		TTGCTGCAAC		_
5960	5970	5980	5990	6000
		CGTCTCGCTG	GTGAAAAGAA	
. 6010	6020	6030	6040	6050
GGCGCCCAAT	ACGCAAACCG	CCTCTCCCCG	CGCGTTGGCC	GATTCATTAA
• 6060	6070	6080	6090	6100
TCCAGCTGGC	ACGACAGGTT	TCCCGACTGG	AAAGCGGGCA	GTGAGCGCAA
6110	6120	6130	6140	6150
CGCAATTAAT	GTGAGTTACC	TCACTCATTA	GGCACCCCAG	GCTTTACACT
6160	6170	6180	6190	6200
		TTGTGTGGAA		
6210	6220	6230	6240	6250
	ACAGCTATGE	CCATGATTAC	GAATTCGLGC	TOGOCOGGG
6260	6270	6280	6290	6300
02.00	02.0	0200	6230	6200

ATCTGCCTGA	ATAGGTACGA	TTTACTAACT		CTAAATGAAC
6310	6320	6330	6340	6350
ACGATTAACA	TCGCTAAGAA	CGACTTCTCT	GACATCGAAC	TGGCTGCTAT
6360	6370	6380	6390	6400
CCCGTTCAAC	ACTCTGGCTG	ACCATTACGG	TGAGCGTTTA	GCTCGCGAAC
6410	6420	6430	6440	6450
AGTTGGCCCT	TGAGCATGAG		TGGGTGAAGC	ACGCTTCCGC
64 60	6470	6480	6490	6500
•	AGCGTCAACT		GAGGTTGCGG	ATAACGCTGC
AAGATGTTTG	6520	6530	6540	6550
6510				GCACGCATCA
CGCCAAGCCT	CTCATCACTA		6590	6600
65 60	6570	6580		CCCGACAGCC
ACGACTGGTT		AAAGCTAAGC	•••	6650
6610	6620	6630	6640	ACATCACCAT
TTCCAGTTCC	TGCAAGAAAT		GCCGTAGCGT	
6660	6670	6680	6690	6700
TAAGACCACT	CTGGCTTGCC	TAACCAGTGC	TGACAATACA	
6710	6720	6730	6740	
CTGTAGCAAG	CGCAATCGGT	CGGGCCATTG	AGGACGAGGC	TCGCTTCGGT
6760	6770	6780	6790	6800
CGTATCCGTG	ACCTTGAAGC	TAAGCACTTC	AAGAAAAACG	TTGAGGAACA
6810	6820	6830	6840	· 6850
	CGCGTAGGGC	ACGTCTACAA	GAAAGCATTT	ATGCAAGTTG
6860	6870	6880	6890	6900
TCGAGGCTGA		AAGGGTCTAC	TCGGTGGCGA	GGCGTGGTCT
_	6920	6930	6940	6950
6910		TATTCATGTA	GGAGTACGCT	GCATCGAGAT
	AGGAAGACTC	6980	6990	7000
6960	6970	• • • •	ACACCGCCAA	AATGCTGGCG
GCTCATTGAG				7050
7010	7020	7030	7040	
TAGTAGGTCA	AGACTCTGAG	ACTATCGAAC	TCGCACCTGA	ATACGCTGAG
7060	7070	7080	7090	7100
GCTATCGCAA	CCCGTGCAGG	TGCGCTGGCT	GGCATCTCTC	CGATGTTCCA
7110	7120	. 7130	7140	7150
ACCTTGCGTA	GTTCCTCCTA	AGCCGTGGAC	TGGCATTACT	GGTGGTGGCT
7160	7170	7180	7190	7200
ATTGGGCTAA	CGGTCGTCGT	CCTCTGGCGC	TGGTGCGTAC	TCACAGTAAG
7210	7220	7230	7240	7250
AAAGCACTGA		AGACGTTTAC	ATGCCTGAGG	TGTACAAAGC
7260	7270	7280	7290	7300
	000011110	CCGCATGGAA		AAAGTCCTAG
GATTAACATT			7340	7350
7310	7320	77000	ATTGTCCGGT	CGAGGACATC
		JOANDO LOAK	7390	7400
7360	7370	7380		
CCTGCGATTG			AAACCOGAAG	ACATEGACAT
· 7410	7420	7430	7440	
GAATCCTGAG	GCTCTCACCG	CGTGGAAACG	TGCTGCCGCT	COLUMBIACO
7460	7470	7480	7490	
GCAAGGACAA	GGCTCGCAAG	TOTOGOCGIA	TCAGCCTTGA	GTTCATGCTT
7510	7520	7530	7540	/550
GAGCAAGCCA	ATAAGTTTGC	TAACCATAAG	GCCATCTGGT	TCCCTTACAA
7560	7570	7580	7590	7603
000. 0070400740	CGCGGTCGTG	TTTACGCTGT	GTCAATGTTC	AACCCGCAAG
7610		7630	7640	7650
7010	, , , ,			

917810011111	GACCAAAGGA		TGGCGAAAGG	TAAACCAATC
7660	7670	7680	7690	7700
	GTTACTACTG			ACTGTGCGGG
7710	7720	7730	7740	7750
TGTCGATAAG	GTTCCGTTCC	CTGAGCGCAT	CAAGTTCATT	GAGGAAAACC
7760	7770	7780	7790	7800
ACGAGAACAT	CATGGCTTGC	GCTAAGTCTC	CACTGGAGAA	CACTTGGTGG
7810	7820	7830	7840	7850
GCTGAGCAAG	ATTCTCCGTT	CTGCTTCCTT	GCGTTCTGCT	TTGAGTACGC
7860	7870	7880	7890	7900
TGGGGTACAG	CACCACGGCC	TGAGCTATAA	CTGCTCCCTT	CCGCTGGCGT
7910	7920	7930	7940	7950
TTGACGGGTC		ATCCAGCACT	TCTCCGCGAT	GCTCCGAGAT
7960	7970	7980	7990	8000
GAGGTAGGTG		TAACTTGCTT	CCTAGTGAAA	CCGTTCAGGA
8010	8020	8030	8040	8050
CATCTACGGG		AGAAAGTCAA		CAAGCAGACG
8060	8070	8080	8090	8100
CAATCAATGG	GACCGATAAC	GAAGTAGTTA	CCGTGACCGA	TGAGAACACT
8110	8120	8130	8140	8150
GGTGAAATCT	CTGAGAAAGT	CAAGCTGGGC	ACTAAGGCAC	TGGCTGGTCA
8160	8170	8180	8190	8200
ATGGCTGGCT	TACGGTGTTA	CTCGCAGTGT	GACTAAGCGT	TCAGTCATGA
8210	8220	8230	8240	8250
CGCTGGCTTA	CGGGTCCAAA	GAGTTCGGCT	TCCGTCAACA	AGTGCTGGAA
8260	8270	8280	8290	8300
GATACCATTC	AGCCAGCTAT	TGATTCCGGC	AAGGGTCTGA	TGTTCACTCA
8310	8320	8330	8340	8350
GCCGAATCAG	GCTGCTGGAT	ACATGGCTAA	GCTGATTTGG	GAATCTGTGA
8360	8370	8380	8390	8400
GCGTGACGGT	GGTAGCTGCG	GTTGAAGCAA	TGAACTGGCT	TAAGTCTGCT
8410	8420	8430	8440	8450
GCTAAGCTGC	TGGCTGCTGA	GGTCAAAGAT	AAGAAGACTG	GAGAGATTCT
8460	8470	. 8480	8490	8500
TCGCAAGCGT	TGCGCTGTGC	ATTGGGTAAC	TCCTGATGGT	TTCCCTGTGT
8510	8520	8530	8540	8550
GGCAGGAATA		ATTCAGACGC	GCTTGAACCT	GATGTTCCTC
8560	8570	8580	8590	8600
GGTCAGTTCC		TACCATTAAC	ACCAACAAAG	ATAGCGAGAT
8610	8620	8630	8.640	8650
TGATGCACAC		CTGGTATCGC	TCCTAACTTT	GTACACAGCC
8660	8670	8680	8690	8700
AAGACGGTAG	CCACCTTCGT		TGTGGGCACA	CGAGAAGTAC
8710	8720	8730		8750
CCARTCAAT	-CTTTTTTCTTC-		TCCTTCGGTA	
8760	8770	8780	8790	8600
			CGAAACTATG	
		8830	8E40	
8810	8820			8950
			ACGACCAGTT 8890	
8860	8870	8880		8900
			GCACTTCCGG	
8910	8920.		8940	8950
			CTTCGCGTTC	
8960	8970	8980	8990	. 9000

FIGURE 9 (continued)

AAATCAATAC	GACCCGGATC	GGTCGACCTG	CAGCCCAAGC	TTGGCACTGG
9010	9020	9030	9040	9050
CCGTCGTTTT	ACAACGTCGT	GACTGGGAAA	ACCCTGGCGT	TACCCAACTT
9060	9070	9080	9090	9100
AATCGCCTTG	CAGCACATCC	CCCCTTCGCC		ATAGCGAAGA
9110	9120	9130	9140	9150
GGCCCGCACC	GATCGCCCTT	CCCAACAGTT	GCGTAGCCTG	AATGGCGAAT
9160	91.70	9180	9190	9200
GGCGCTTTGC	CTGGTTTCCG	GCACCAGAAG	CGGTGCCGGA	AAGCTGGCTG
9210	9220	9230	9240	9250
GAGTGCGATC	TTCCTGAGGC		GTCGTCCCCT	CAAACTGGCA
9260	9270	9280	9290	9300
GATGCACGGT		CCATCTACAC	CAACGTAACC	TATCCCATTA
9310	9320	9330	9340	9350
CGGTCAATCC		CCCACGGAGA		TTGTTACTCG
9360	9370	9380	9390	9400
CTCACATTTA		AAGCTGGCTA		AGACGCGAAT
9410	9420	9430	9440	9450
TATTTTTGAT		TTGGTTAAAA	AATGAGCTGA	TTTAACAAAA 9500
9460	9470	9480	9490	TAAATATTTG
ATTTAACGCG		AAATATTAAC 9530	9540	9550
9510	9520		TCTGATTATC	AACCGGGGTA
CTTATACAAT	CTTCCTGTTT	9580	9590	9600
9560	9570	• • • •	CCGTTCATCG	ATTCTCTTGT
CATATGATTG	ACATGCTAGT 9620	9630	9640	9650
9610			AGCCTTTGTA	GATCTCTCAA
TTGCTCCAGA	CTCTCAGGCA 9670	9680	9690	9700
AAATAGCTAC	CCTCTCCGGC	ATGAATTTAT	CAGCTAGAAC	
9710	9720	9730	9740	9750
CATATTGATG	GTGATTTGAC	TGTCTCCGGC		CTTTTGAATC
9760	9770	9780	9790	9800
TTTACCTACA	CATTACTCAG	GCATTGCATT	• • • -	GAGGGTTCTA
9810	9820	9830	9840	9850
AAAATTTTTA		GAAATAAAGG	• • • •	
9860	9870	9880	9890	9900
CAGGGTCATA	ATGTTTTTGG	TACAACCGAT		GCTCTGAGGC
9910	9920	9930	9940	9950
TTTATTGCTT		ATTCTTTGCC		GATTTATTGG

ATGTT

FIGURE 10

